THORPE'S DICTIONARY OF APPLIED CHEMISTRY

BY

(the late)

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ASSISTED BY EMINENT CONTRIBUTORS

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${ t FOREWORD}$

VOLUME III has been constructed on the same lines as those followed in Volume II. The Foreword written in Volume II therefore covers all that it is necessary to say

concerning Volume III.

Dr. Johnson says in the Preface to his Dictionary, published in 1755, "It is the fatc of those who toil at the lower employments of life to be rather driven by the fear of evil, than attracted by the prospect of good; to be exposed to censure, without hope of praise; to be disgraced by miscarriage, or punished for neglect, where success would have been without applause, and diligence without reward.

"Among these unhappy mortals is the writer of dictionaries; whom mankind have

considered, not as the pupil, but the slave of science, the pionier of literature, doomed only to remove rubbish and clear obstructions from the path of Learning and Genius, who press forward to conquest and glory, without bestowing a smile on the humble drudge that facilitates their progress. Every other author may aspire to praise; the lexicographer ean only hope to escape reproach, and even this negative recompense has been yet granted to very few.

"... This recommendation of steadiness and uniformity does not proceed from an opinion, that particular combinations of letters have much influence on human happiness; or that truth may not be successfully taught by modes of spelling fanciful and erroneous: I am not yet so lost in lexicography, as to forget that 'words are the daughters of earth, and that things are sons of heaven.' Language is only the instrument of science, and words are but the signs of ideas: I wish, however, that the instrument might be less apt to decay,

and that signs might be permanent, like the things which they denote.

". . . To deliberate whenever I doubted, to enquire whenever I was ignorant, would have protracted the undertaking without end, and, perhaps, without much improvement; for I did not find by my first experiments, that what I had not of my own was easily to be obtained: I saw that one enquiry only gave occasion to another, that book referred to book, that to search was not always to find, and to find was not always to be informed; and that thus to persue perfection, was, like the first inhabitants of Arcadia, to chace the sun, which, when they had reached the hill where he seemed to rest, was still beheld at the

same distance from them.

". . . I look with pleasure on my book, however defective, and deliver it to the world with the spirit of a man that has endeavoured well. That it will immediately become popular I have not promised to myself: a few wild blunders, and risible absurdities, from which no work of such multiplicity was ever free, may for a time furnish folly with laughter, and harden ignorance in contempt; but useful diligence will at last prevail, and there never can be wanting some who distinguish desert; who will consider that no dictionary of a living tongue ever can be perfect \dots even a whole life would not be sufficient; that he, whose design includes whatever language can express, must often speak of what he does not understand; that a writer will sometimes be hurried by eagerness to the end, and sometimes faint with weariness under a task, which Scaliger compares to the labours of the anvil and the mine; that what is obvious is not always known, and what is known is not always present; that sudden fits of inadvertency will surprize vigilance, slight avocations will seduce attention, and casual eclipses of the mind will darken learning; and that the writer shall often in vain trace his memory at the moment of need, for that which yesterday he knew with intuitive readiness, and which will come uncalled into his thoughts to-morrow."

There can be no doubt that the great man had his tongue in his cheek when he

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ABBREVIATIONS

OF THE TITLES OF JOURNALS, ETC.

(Reproduced by permission of the Bureau of Abstracts.)

A. I; A. II; A. III	British Chemical and Physiological Abstracts.
Abh. Böhm. Akad	Abhandlungen der Bohmischen Akadeinie.
Acta Bot. Fenniea	Acta Botanica Fennica.
Acta Physicochim. U.R.S.S.	Acta Physicochimica U.R S.S.
Acta Phytochim	Acta Phytochimica.
Acta Sci. Fennica	Acta Societatis Scientiarum Fennicæ.
Agric. Eng	Agricultural Engineering.
Agric. Gaz. New South Wales	
Agric, J. Brit. Guiana	Agricultural Journal of British Guiana.
Agric. Live-stock India	Agriculture and Live-stock in India.
Agrie. Res. Inst., Pusa, Rep.	
(Bull.)	Agricultural Research Institute, Pusa, Reports and Bulletins.
Allgem. Ocl- Fett-Ztg	Allgemeine Oel- und Fett-Zeitung.
Alloem Z Rierbrau	Allgemeine Zeitschrift für Bierbrauerei und Malzfabrikation.
.1luminium	Aluminum. Chemical Abstracts. Published by the American Chemical Society. American Dyestuff Reporter. American Gas Journal.
Amer. Chem. Abstr	Chemical Abstracts. Published by the American Chemical Society.
Amer. Duestuff Ren	American Dyestuff Reporter.
Amer. Gas J	American Gas Journal.
Amer. Ink Maker	American Ink Maker.
Amer. Inst. Min. Met. Eng.	
Publ.	tion.
Amer. J. Bot	American Journal of Botany.
Amer. J. Dis. Children	American Journal of Diseases of Children.
Amer. J. Pharm	American Journal of Pharmacy.
Amer. J. Physiol	American Journal of Physiology.
Amer. J. Publ. Health	American Journal of Public Health and the Nation's Health.
Amer. J. Sci	American Journal of Science.
Amer. Min	American Mineralogist.
Amer. Paint J	American Paint Journal.
Amer. Potato J	American Potato Journal.
	Anaics de la Asociación Química Argentina.
Anal. Fis. Quim.	Anales de la Sociedad Española de Física y Quimica.
	Anales del Instituto de investigaciones científicas y tecnologicas.
Analyst	Analyst.
Anat Rec	Anatomical Record.
Angew. Bot	Angewandte Botanik.
	Angewandto Chemie (changed from Z. angew. Chem. in 1932).
Anılınokras. Prom	Androkrasotschnaja Promischlennosti.
	Annals da Academia Brasileira de Sciencias.
Annalen	Annales Academie Scientiarum Fennice.
	Justus Liebig's Annalca der Chemie.
Ann. agron	Annales agronomique.
Ann Rot	Annals of Retary
Annalı Chim Annl	Aungle de Chimage Applicate
Ann. Chim.	Annales de Chimie
Ann Chim, Analut.	Arnales de Chimie Analytique et de Chimie Analysis
Ann. di Bot.	Annales agronomique. Annales of Applied Biology. Annale of Botany. Annale di Chimica Applicata. Annales de Chimic. Annales de Chimic Analytique et de Chimic Appliquée. Annale di Botanica. Annales des Falsifications.
Ann. Falsif	Annales des Falsifications
Ann. Ferm.	Annales des Fermentations
Ann Inst. Anal. Phys. Chim.	Annales des Fermentations. Annales de l'Institut d'Analyse Physico-chimique.
Ann. Inst. Pasteur	Annales de l'Institut Pasteur.
Ann. Mines Belg	Annales des Mines de Belgique.
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BAL Uff. Staz. Sperim. Ind. R. Stazione Sperimentale per l'Industria delle Pelli e delle materie
                                  concianti, Bollettino Ûfficiale.
  Pelli.
Bot. Archiv . .
                             Botanisches Archiv.
Bot. Gaz. .
                            Botanical Gazette.
Brass. Malt. .
                          . Brasserie et Malterie.
                          . Brau- u. Malzindustrie.
Brau- u. Malzind. .
                          . Braunkohle.
Braunkohle . .
Braunkohlenarchiv .
                             Braunkohlenarchiv.
Brennetoff-Chem.
                             Brennstoff-Chemie.
                          . Brewers' Journal.
Brewers' J.
Brewers' J. . . Brit. Dental J. .
                            British Dental Journal.
Brit. Guiana Dept. Agric.
                             British Guiana Department of Agriculture Bulletin.
  Bull.
                             British Journal of Experimental Pathology.
Brit, J. Exp. Path. . .
                             British Journal of Photography.
Brit, J. Phot.
Brit. J. Phys. Med. .
                             British Journal of Physical Medicine.
Brit. Med. J.
                             British Medical Journal.
                             British Plastics and Moulded Products Trader.
Brit. Plastics .
Bul. Chim.
                             Buletinul Chimie.
Bul. Chim. Soc. Române
                             Buletinul de Chimie pura si aplicata al Societatii Româue de Chimie.
Bul. Soc. Chim. România .
                             Buletinul Societății de Chimie din România.
Bul. Soc. Fiz. România .
                             Buletinul Bilunar al Societății de Fizică din România.
Bul. Soc. Române Stiin.
                             Buletinul Societatii Române de Stiinte.
Bull. Acad. Méd. Roumanie
Bull. Acad. Polonaiss
                             Bulletin de l'Académie de Médecine de Roumanie.
                             Bulletin Internationale de l'Académie Polonaise des Sciences et des
                                  Lettres.
                             Académic royale de Belgique—Bulletin de la Classe des Sciences.
Bull. Acad. rov. Belg.
Bull. Acad. Sci. Agra &
                             Bulletin of the Academy of Sciences of Agra and Oudh.
Bull. Acad. Sci. Roumaine .
                             Bulletin de la Section Scientifique de l'Académie Roumaine.
Bull. Acad. Sci. U.R.S.S.
                             Bulletin de l'Académie des Sciences de l'Union des Républiques
                                  Soviétiques Socialistes.
Bull. Agric. Chem. Soc.
Bulletin of the Agricultural Chemical Society of Japan.
                             Bulletin of the American Ceramic Society.
Bull. Assoc. Chim. Sucr.
                             Bulletin de l'Association des Chimistes de Sucrerie et de Distillerie.
Bull. Biol. Méd. exp.
  U.R.S.S.
                             Bulletin de Biologie et Médecine expérimentale de l'U.R.S.S.
Bull. Chem. Soc. Japan
                             Bulletin of the Chemical Society of Japan.
Bull. Dept. Agric. Kenya
                             Bulletin of the Department of Agriculture of Kenya.
Bull. Dept. Agric. South
  Africa .
                             Bulletin of the Department of Agriculture of South Africa.
Bulletin of Entomological Research.
  Meguro
                             Bulletin of the Forest Experiment Station, Meguro, Tokyo.
Bull. Imp. Inst. .
                             Bulletin of the Imperial Institute.
Bull. Inst. Min. Met.
                             Bulletin of the Institution of Mining and Metallurgy.
Bull. Inst. Phys. Chem. Res.
                             Bulletin of the Institute of Physical and Chemical Research, Japan
  Japan.
                                 (Rikagaku Kenkyujo Ihō).
Bull. Inst. Pin
                             Bulletin de l'Institut du Pin.
Bull. Johns Hopkins Hosp.
                             Bulletin of the Johns Hopkins Hospital.
Bull. Mat. Grasses .
                             Bulletin des Matières Grasses.
Bull. Photogrammétrie
                             Bulletin de Photogrammétrie.
Bull. Rubber Growers' Assoc.
                             Bulletin of the Rubber Growers' Association.
Bull. School Mines and Met.,
                             Bulletin of the School of Mines and Metallurgy, University of
  Unic. Missouri.
Bull. Sei. Pharmacol.
                             Bulletin des Sciences Pharmacologiques.
Bull. Sericult. Japan ... Bull. Soc. chirt. . . . .
                             Bulletin of Sericulture and Silk Industry, Japan.
                             Bulletin de la Société chimique de France.
Bull. Soc. chim. Belg.
                             Bulletin de la Société chimique de Belgique.
Bull. Soc. Chim. biol.
                             Bulletin de la Société de Chimie biologique.
Bull. Soc. Chim. Yougoslav.
                             Bulletin de la Société Chimique du Royaume de Yougoslavie,
Bull. Soc. & Encour. . .
                             Bulletin de la Société d'Encouragement pour l'Industrie Nationale.
Bull. Soc. Franc. Min. .
                             Bulletin de la Société Française de Minéralogie.
Bull. Soc. France. Phot. .
                          . Bulletin de la Société Française de Photographie et de Cinémato-
                                 graphie.
Bull. Sec. Ind. Mulhouse.

Bull. Sec. Sci. Hyg. aliment.

Bull. U.S. Geel. Survey.

Bulletin of the U.S. Geological Survey.

Canala Department of Mines Publications.
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Initanda	10,40, 04 316. 111 01 0001
Empire Colton Growing Rev.	Empire Cotton Growing Review.
Empire J. Exp. Agric	Empire Journal of Experimental Agriculture.
Engineer	Engineer.
Engineering	T
	1 22 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
	Enzymologia.
Enzymologia	
Ergebn. Physiol	Distriction of the state of the
Ernährung	Die Ernährung.
Ernähr. Pflanze	Ernährung der Pflanze.
F.P	French Patent.
Facts about Sugar	Laets about Sugar.
Farbe u. Lack	Farbe und Lack.
Farben-Chem	Farben-Chemiker.
Farben-Ztg	Farben-Zeitung.
Farve og Lak Fermentforsch	Farve og Lak.
Fermenlforsch	Fermentforschung.
Fert. Feeding Stuffs J	Fertiliser, Feeding Stuffs and Farm Supplies Journal.
Felle u. Seifen	Fette und Seifen.
Feuerfest	Feuerfest.
Feuerungsteeh	1 =
Pinaha Pan Madd	Finska Kemistsamfundets Meddelanden (Suomen Kemistiseuran
Finska Kein. Medd	l = · · · · · · · · · · · · · · · · ·
F-1	Tiedonantoja).
Flora	Flora.
Florida Agric. Exp. Sta.	
Bull	Florida Agricultural Experiment Station Bulletin.
Food	Food.
Food Manuf	Food Manufacture.
Food Res	Food Research.
Forstarchiv	Forstarchiv.
Foundry Trade J	Foundry Trade Journal.
Fruit Prod. J	Fruit Products Journal.
Fuel	Fuel in Science and Practice.
Fuel Econ	Fuel in Science and Practice. Fuel Economist.
Fuel L'on. Rev	Fuel Feenamy Derion
Fuel L'on. Rev	Fuel Economy Roview. German Patent.
Q.P.	German Patent.
Gas Ind	Gas Industry.
Gas J.	Gas Journal.
Gas Times	Gas Times.
Gas- u. Wasserfach	Gas- und Wasserfach.
Gas World	Gas World.
Gazzella	Gazzetta chimica italiana.
Geol. Mag	Geological Magazine.
Georgia Agric. Exp. Sta.	
Bull.	Georgia Agricultural Experiment Station Bulletin.
Ges. Abh. Kennt. Kohle	Gesammelto Abhandlungen der Kenntnis der Kohle.
Glass Ind.	Glass Industry.
Classicus	
Glasshütte	Die Glasshütte.
Glückauf	
Gummi-Ztg	Gummi-Zeitung.
Hawaii Agric. Exp. Stat.	77 11 4 12 14 1 77 1 1 1 100 100 100 100 100
Bull.	Hawaii Agricultural Experiment Station Bulletins.
Helv. Chim. Acta	Helvetica Chimica Acta.
Hilgardia	Hilgardia.
Imp. Bur. Soil Sci. Tech.	
Comm	Imperial Bureau of Soil Science, Technical Communications.
Indian J. Agric. Sci	Indian Journal of Agricultural Science.
Indian J. Med. Res	Indian Journal of Medical Research.
Indian J. Physics	Indian Journal of Physics.
Indian J. Vet. Sei	Indian Journal of Veterinary Science.
Indian Lac Res. Inst. Bull.	Indian Lac Research Instituto Bulletin.
India-rubber J.	India-rubber Journal.
Ind. Chem.	Industrial Chemist.
Ind. Eng. Chem.	
Ind. Eng. Chem. [Anal.]	Industrial and Engineering Chemistry.
Int Day 1	Industrial and Engineering Chemistry, Analytical Edition.
Int. Sugar T	International Review of Agriculture.
Int. Sugar J	International Sugar Journal.
Iron Age	Iron Age.
Iron Steel Inst. Carnegie Schol. Mem.	
ACROL Mem	
100	Iron and Steel Institute, Carnegie Scolarship Memoirs.
J.C.S.	Iron and Steel Institute, Carnegie Scolarship Memoirs. Journal of the Chemical Society.

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J. Fuel Soc. Japan .
                             Journal of the Fuel Society of Japan.
                             Journal of General Chemistry, Russia (formerly J. Russ. Phys.
J. Gen. Chem. Russ.
                                  Chem. Soc.).
                             Journal of General Physiology.
J. Gen. Physiol. . .
                             Journal of Geology.
J. Geol.
                             Chishitsugaku Zasshi (Journal of the Goological Society of Tokyo).
J. Geol. Soc. Tokyo .
J. Hygiene . .
                             Journal of Hygiene.
J. Indian Chem. Soc.
                             Journal of the Indian Chemical Society.
                             Journal of the Indian Institute of Science.
J. Indian Inst. Sci. .
                             Journal of Industrial Hygicne and Toxicology.
J. Ind. Ilyg. . .
J. Infect. Dis.
                             Journal of Infectious Diseases.
J. Inst. Brew.
                             Journal of the Institute of Brewing.
                             Journal of the Institution of Electrical Engineers.
J. Inst. Electr. Eng. .
                             Journal of the Instituto of Fuel.
J. Inst. Fuel.
J. Inst. Metals .
                             Journal of the Institute of Metals.
J. Inst. Petroleum Tech.
                             Journal of the Institution of Petroleum Technologists.
                             Journal of the Institute of Sewage Purification.
Journal of the Iron and Steel Institute.
J. Inst. Sewage Purif.
J. Iron and Steel Inst.
                             Journal of the Japanese Ceramic Association.
J. Jap. Ccram. Assoc.
J. Landw.
                             Journal für Landwirtschaft.
J. Marine Biol. Assoc. .
                             Journal of the Marine Biological Association of the United Kingdom.
J. Med. Res. . . .
                             Journal of Medical Research.
                             Journal of the Ministry of Agriculture.
J. Min. Agric.
J. Min. Agric. N. Ireland
                             Journal of the Ministry of Agriculture of Northern Ireland.
    New England Water
  Works Assoc. . .
                             Journal of the New England Water Works Association.
                             Journal of Nutrition.
J. Nutrition .
J. Oil Col. Chem. Assoc.
                             Journal of the Oil and Colour Chemists' Association.
J. Opt. Soc. Amer. . .
                             Journal of the Optical Society of America.
J. Org. Chem.
                             Journal of Organic Chemistry.
                             Journal of Pathology and Bacteriology.
J. Path. Bact.
J. Pharm. Chim.
                             Journal de Pharmacic et de Chimie.
J. Pharm. Exp. Ther.
                             Journal of Pharmacology and Experimental Therapeutics.
J. Pharm. Soc. Japan
                             Journal of the Pharmaccutical Society of Japan. (Yakugakuzasshi.)
J. Physical Chem. .
                             Journal of Physical Chemistry.
J. Phys. Chem. Russ.
                              Shurnal Fizitscheskoi Chimii.
                             Journal of Physiology.
J. Physiol.
J. Physiol. Path. gén.
                              Journal de Physiologie et de Pathologie généralc.
J. Phys. Radium .
                             Journal de Physique et le Radium.
J. Pomology .
                             Journal of Pomology and Horticultural Science.
J. pr. Chem. .
                             Journal für praktische Chemie.
J. Proc. Asiatic Soc. Bengal
                             Journal and Proceedings of the Asiatic Society of Bengal.
J. Proc. Austral. Chem. Inst.
                             Journal and Proceedings of the Australian Chemical Institute.
J. Proc. Roy. Soc. New South
                             Journal and Proceedings of the Royal Society of New South Wales.
J. Res. Nat. Bur. Stand.
                             Journal of Research of the National Bureau of Standards.
J. Roy. Agric. Soc. . .
                             Journal of the Royal Agricultural Society.
J. Roy. Hort. Soc.
                             Journal of the Royal Horticultural Society.
J. Roy. Microscop. Soc.
                             Journal of the Royal Microscopical Society.
J. Roy. Soc. West Australia
                             Journal of the Royal Society of West Australia.
J. Roy. Tech. Coll. .
                             Journal of the Royal Technical College (Glasgow).
J. Rubber Res.
J. Rubber Res. Inst. Malaya
                             Journal of Rubber Research.
                             Journal of the Rubber Research Institute of Malaya.
J. Sci. Hiroshima Univ.
                              Journal of Science of the Hiroshima University.
                             Journal of Scientific Instruments.
Journal of Science and Technology, India.
J. Sci. Instr.
J. Sci. Tech. India .
                             Journal of the Scottish Meteorological Society.
J. Scot. Mct. Soc.
J. Soc. Arts . .
                             Journal of the Royal Society of Arts.
J.S.C.I.
                             Journal of the Society of Chemical Industry.
J. Soc. Chem. Ind. Japan
                             Journal of the Society of Chemical Industry, Japan. (Kögyö
                                  Kwagaku Zasshi.)
J. Soc. Dyers and Col. .
                             Journal of the Society of Dyers and Colourists.
J. Soc. Glass Tech. .
                             Journal of the Society of Glass Technology.
J. Soc. Leather Trades'
  Chem.
                             Journal of the International Society of Leather Trades' Chemists.
J. S. African Chem. Inst.
                             Journal of the South African Chemical Institute.
J. South-East. Agric. Coll. .
                             Journal of the South-Eastern Agricultural College, Wye, Kent.
J. State Med.
                             Journal of State Medicine.
J. Text. Inst.
                             Journal of the Textile Institute.
J. Univ. Bombay
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J. Usines Gaz	Journal des Usmes à Gaz.
J. Washington Acad Sci .	Journal of the Washington Academy of Sciences.
J. Danington Acad Des .	
J. West Scotland Iron Steel	The state of the s
Inst	Journal of the West of Scotland Iron and Steel Institute.
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Last	Le Last.
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Montana Agric. Exp. Sta.	
Bull.	Montana Agricultural Experiment Station Bulletin.
Month. J. Inst. Metals	Monthly Journal of the Institute of Metals.
Month. Not. Roy. Astr. Soc.	Monthly Notices of the Royal Astronomical Society, London.
Month. Rep. Dept. Agric. N.	Monthly Reports of the Department of Agriculture of Northern
Ireland.	Ircland.
Mühlenlab	Das Mühlenlaboratorium.
Münch. med. Woch	Münchener medizinische Woehensehrift.
Numer. med. 11 ben	Nachrichten von der Geschlschaft der Wissensehaften zu Göttingen.
	National Butter and Cheese Journal.
Nat. Butter & Cheese J.	National Daint Marich and Laguer Association Inc. Circular
Nat. Paint Var. Assoc. Circ.	National Paint, Varnish, and Lacquer Association, Inc., Circular.
Nature	Nature.
Natuurwetensch. Tijds	Natuurwetenschappelijk Tijdschrift.
Naturwiss	Die Naturwissenschaften.
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Sta. Bull	New Hampshire Agricultural Experiment Station Bulletin.
New Jersey Agric. Exp. Sta.	
Bull	Now Jersey Agricultural Experiment Station Bulletin.
New Phytol	New Phytologist.
New York Agr. Expt. Sta.	
Bull	Now York State Agricultural Experiment Station Bulletins.
New York (Geneva) Agric.	
Exp. Sta. Bull	Now York (Geneva) Agricultural Experiment Station Bulletin.
New Zealand Dominion	
Laby. Rept	New Zealand Dominion Laboratory Reports.
New Zealand J. Sci. Tech	New Zcaland Journal of Science and Technology.
Nova Acta Soc. Sci	Nova Acta Regiae Societatis Scientiarum Upsaliensis.
Nutr. Abs	Nutrition Abstracts and Roviews.
Ofvers. Finska Vet. Soc	Öfversigt af Finska Vetenskaps-Societétens Förhandlingar,
•	Helsingfors.
Oel u. Kohle	Ocl und Kohle.
Off. Digest	Official Digest of the Federation of Paint and Varnish Production
	Clubs.
Ohio Agric. Exp. Sta. Bull.	Ohio Agricultural Experiment Station Bulletin.
Oil and Gas J	Oil and Gas Journal.
Oil and Soap	Oil and Soap.
Oklahoma Agric. Exp. Sta.	_
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Pacific Pulp and Paper Ind.	Forhandlingar.
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Physikal. Z.
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                             Phytopathology.
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                             Proceedings of the Academy of Sciences of the United Provinces of
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                                  tion of Great Britam and Ireland
Proc. U.S. Nat Mus
                             Proceedings of the United States National Museum.
Protoplasma .
                             Protoplasma
Przemyst Chem
                             Przemysł Chemiczny
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                             Publications de la Faculte des Sciences de l'Université Masaryl.
  Masaryk
                                  (Spisy vydavané Prírodovedeckou Pacoultou Masarykovy
                                  University)
Pulp and Paper
                      Mag
                             Pnlp and Paper Magazine of Canada
  Canada
Quart. J. Exp. Physiol
                             Quarterly Journal of Experimental Physiology.
                             Quarterly Journal of the Geological Society.
Quart. J. Geol Soc
Quart. J. Med
                             Quarterly Journal of Medicine
Quart. J. Pharm
                             Quarterly Journal of Pharmacy and Pharmacology
Queensland J. Agrıc
                             Queensland Journal of Agriculture.
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Rayon Textile Monthly

Records of the Australian Museum

Recueil des travaux bottinques Acerlandaises Recueil des travaux chimiques des Pays Bas

ABBREVIAL	IONS OF THE TITLES OF JOURNALS, ETC. XXI
Refiner	Refiner and Natural Gasolinc Manufacturer. Rendiconto dell' Accademia delle Scienze Fisiche e Matematiche, Napoli.
Napoli	Rendiconti doll' Reale Istituto Lombardo di Scienze e Lettere. Rensselaer Polytechnic Institute Bulletin.
Rep. Aust. Assoc. Sci	Report of the Australian Association for the Advancement of Science.
Rep. Brit. Assoc	Roport of the British Association for the Advancement of Science.
ington	Research Studies of the State College of Washington. Rovue de l'Aluminium.
Rev. Chim. pura appl	Revista de Chimica pura e applicada, Orgão de Sociedade Portuguêsa do Química e Física.
Rev. Fac. Quim. Ind. Agric.	Revista de la Facultad de Quimica Industrial y Agrícola.
Rev. Gén. Mat. Col	Revue Générale des Matières Colorantes. Revuo do Métallurgie.
Rev. Mod. Physics	Review of Modern Physics.
Rev. Sci. Instr	Review of Scientific Instruments.
Rocz. Chem	Roczniki Chemji organ Polskiego Towarzystwa Chemicznego.
Shorn. Trud. Ukrain. Chim. Inst. Odcssa	Sbornik Trudov Ukrainskogo Nautschno-Issledova-Telskogo Chimitscheskogo Instituta Odessa.
Science	Science.
Sci. Agric	Scientific Agriculture. Science and Culture.
Sci. and Cult	Science et Industric.
Sci. et Ind	Dolondo of Industrio.
Bertrand Fils	Scientific and Industrial Roports of Roure-Bertrand Fils.
Sci. Mcm. Univ. Saratov .	Scientific Memoirs of the University of Saratov.
Sci. Papers Inst. Phys. Chem.	Scientific Papers of the Institute of Physical and Chemical Research,
Res. Tokyo	Tokyo.
Sci. Proc. Roy. Dublin Soc.	Scientific Proceedings of the Royal Dublin Society.
Sci. Quart. Nat. Univ. Peking Sci. Rep. Hiroshima Tech.	Science Quarterly of the National University of Peking.
Sch	Scientific Reports of the Hiroshima Higher Technical School.
Sci. Rep. Tohoku	Science Reports, Tolloku Imperial University.
Sci. Rep. Tsing Hua Univ.	Science Reports of the National Tsing Hua University.
Sci. Sect. Nat. Paint, Var.	
Assoc. Circ.	Inc., Circulars.
Sci. Trans. Roy. Dubl. Soc. Scot. J. Agric.	Scientific Transactions of the Royal Dublin Society. Scottish Journal of Agriculture.
ScifensZtg	Seifensieder-Zeitung.
Sewage Works J	Sewage Works Journal.
Sitzungsber. Akad. Wiss.	
Wien	Sitzungsberichte der Akademie der Wissenschaften, Wien.
Sitzungsber. Heidelberger Akad. Wiss	Sitzungsberichte der Heidolberger Akademie der Wissenschaften.
Sitzungsber. Preuss. Akad. Wiss. Berlin	Sitzungsberichte der Preussischen Akademie der Wissenschaften zu Berlin.
Smithsonian Misc. Coll	Smithsonian Miscellaneous Collection.
Soap	Soap.
Soap Trade Rev.	Soap, Perfumery and Cosmetic Trade Review.
Soil Res	Soil Research.
Sparwirts.	Soil Science.
Sprechsanl	Sparwirtschaft. Sprechsaal.
Stahl u. Eisen	Stahl und Eisen.
Stain Tech	Stain Technology.
Staz. sper. agr. ital	Stazioni sperimentali agranic italiane.
Steam Enq	Steam Engineer.
Sugar Bull	Sugar Bulletin.
Superphosphat	Suomen Kemistilehti Acta Chemica Fennica. Superphosphat.
Superphosphate	Superphosphate.
Stensk Kem, Tidskr.	Svensk Kemisk Tidskrift.
Tasmanian Agric. J	Tasmanian Agricultural Journal.
Tech. Mitt. Krupp	Technische Mitteilungen Krupp.
Tech. Publ. Tin Res. Counc.	Technical Publications of the International Tin Research and
Tech. Rep. Tohoku	Development Council. Technology Reports of the Tohoku Imperial University, Sendai,

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XXII
                              Teer and Bitnmen.
Teer u Buumen .
                              Tekni-k Tidakrift.
Tekn. Tidsk. .
                              Melband Textilbenchte.
Textilber. .
                              Tidsskrift for Kjemi og Bergvesen.
Tida Kiemi .
Tonind . Zig. ,
                              Tonindustrie-Zeitung.
Trans. Amer. Inst. Chem.
                              Transactions of the American Institute of Chemical Engineers.
                              Transactions of the American Institution of Metals.
 Trans. Amer. Inst. Metals .
                              Transactions of the American Institute of Mining and Metallurgical
Trans. Amer Inst Min
   Met. Eng. .
                                   Engineers
Trans. Amer. Soc. Mech
                              Transactions of the American Society of Mechanical Engineers.
   Eng
                              Transactions of the American Society of Metals.
Trans. Amer. Soc. Met.
                              Transactions of the Ceramic Society
Trans. Ceram. Soc.
                              Transzerions of the Doknchaiev Soil Institute.
Trans. Dokuchasev Soil Inst.
                              Transactions of the Electrochemical Society,
Trans Lleetrochem Soc.
Trans. Faraday Soc
                              Transactions of the Faraday Society.
Trans, Inst Min Eng.
                              Transactions of the Institution of Mining Engineers
Trans Inst Min. Met
                              Transactions of the Institution of Mining and Metallurgy.
                              Transactions of the Institute of the Plastics Industry,
Trans Inst Plast Ind
                              Tran-actions of the In-titution of the Rubber Industry
Trans Inst Rubber Ind.
                              Transactions of the North of England Institute of Mining and
Trans A Eng. Inst. Min
   Mech Eng
                                   Mechanical Engineers
                              Transactions of the New Zealand Institute.
Transactions of the Nova Scotia Institute of Science.
Trans New Zealand Inst
Trans Nova Scotia Inst Sea
Trans. Opt Soc
                              Transactions of the Optical Society
                              Transactions of the Royal Irish Academy
Trans Roy Irish Acad
Trans. Roy. Soc Canada
                              Transactions of the Royal Society of Canada
Trans Roy Soc Edin
                              Transactions of the Royal Society of Edinburgh.
Trans Roy Soc S Africa
                              Transactions of the Royal Society of South Africa
Trans Saratov Unit
                              Transactions of the Saratov University (Gelehrte Notizen der
                                   Saratover Staats Universität).
Trop. Agric .
                              Tropical Agriculturist (Ceylon).
Trop Agric. (Trinidad)
Tech. Min Miti
                              Tropical Agriculture (Trinidad)
Mineralogische und Petrographische Mitteilungen (Zeitschrift für
                              Kristallographie, Mineralogie, und Petrographie, Abteilung B)
United States Burean of Mines, Bulletins, Technical Papers, and
       Bur. Mines, Bull .
   Tech Papere, and Rept
                                  Reports of Investigations.
  Invest.
U.S. Bur Plant Ind.
                              United States Bureau of Plant Industry,
U.S Dept. Agric. Bull. (or
  Circ ) .
                              Umited States Department of Agriculture Bulletins (or Circulars).
U.S. Hy2 Labor, Bull
                              Umted States Hygiemic Laboratory Bulletins,
U.S.P.
                              United States Patent.
U.S. Publ Health Rep .
                              United States Public Health Reports.
Ukrain Biochem J.
                              Ukraman Biochemical Journal.
Ukrain Chem. J
                              Ukrainian Chemical Journal
Union S Africa Dept Agric.
                              Umon of South Africa Department of Agriculture Bulletins.
  Bulls
Univ. Illinois Bull .
                              University of Illinois Bulletins.
Utah Agric. Coll Exp Stat
  Bull
                              Utah Agricultural College Experiment Station Bulletins.
l'erfkroniel.
                              Verfkromek.
Verh Geol Reschwanst Wien
                             Verhandlungen der Geologischen Reichsanstalt in Wien.
Verh. Ges deut. Naturforsch
                             Verhandlungen der Gesell-chaft deutscher Naturforscher und
  Aertze,
                                  Acrtze.
l'ermont Agric. Exp. Sta.
  Bull.
                             Vermont Agricultural Experiment Station Bulletin.
Vet. Rec
                             Veternary Record.
Virginia (Blacksburg) Exp
  Sta. Bull
                             Virginia (Blacksburg) Experiment Station Bulletin.
Virginia Truck Exp. Sta
  Bull
                             Virginia Truck Experiment Station Bulletin.
Welsh J. Agric.
                             Welsh Journal of Agriculture.
W 145
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                   Stemens
  Werken
                             Wissenschaftliche Veröffentlichungen aus den Siemens-Werken.
Woch Brau.
                             Wochenschrift für Brauerei
World's Paper Tr. Rer .
                             World's Paper Trade Review.
Z. anal Chem.
                             Zeitschrift fnr analytische Chemie.
Z. anorg. Chem
                            Zeitschrift für anorganische und allgemeine Chemie.
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Z. Biol	Zeitschrift für Biologie.
Z. deul. Geol. Ges	Zeitschrift der deutschen Geologischen Gesellschaft.
Z. Elel.trochem	AT
Z. Fleisch- Milch-Hyg	Zeitschrift für Fleisch- und Milch-Hygiene.
Z. ges. Brauv.	Zeitschrift fur das gesamte Brauwesen.
Z. ges. exp. Med	Zeitschrift fur die gesamte experimentelle Medizin.
Z. ges. Naturwiss.	Zeitschrift für die gesamte Naturwissenschaft.
Z. ges. Schiess- u. Spreng-	201130d1119 141 the Bostinte Minister Macondonies
	Zeitschrift fur das gesamte Schiess- und Sprengstoffwesen.
stoffw	Zeitschrift fur Hygiene und Infektionskrankheiten.
Z. Hyg	Zeitschrift für Instrumentenkunde.
Z. Instrumkde	
Z. Krist Z. Mctallk	Zeitschrift für Metallkunde.
	Zeitschrift für Parasitenkunde.
Z. Parasitenk	Zeitschrüt für Parasitenkunde.
Z. Pflanz. Düng	Zeitschrift fur Pflanzanernährung, Düngung, und Bedenkunde
7 DA 1 DA	(title now changed to Bedenkunde und Pflanzenernahrung).
Z. Pflanzenkr. Pflanzen-	
schutz.	Pflanzenschutz.
Z. Physik	Zeitschrift fur Physik.
Z. physikal. Chem	Zeitschrift für physikalische Chemic.
Z. physikal. chem. Unterr	Zeitschrift für den physikalischen und chemischen Unterrieht.
Z. physiol. Chem	Heppe Seyler's Zeitschrift fur physiologische Chemie.
	Zeitschrift fur praktische Geologie.
Z. Spiritusind	Zeitschrift fur Spiritusindustrie.
Z. Unters. Lebensm	Zeitschrift fur Untersuchung der Lebensmittel.
Z. Ver. deul. Ing Z. Ver. deul. Zucker-Ind	Zeitschrift Vereins deutscher Ingenieure.
Z. Ver. deut. Zucker-Ind	
	changed to Z. Wirts. Zuckerind.).
Z. Vitaminforsch	Zeitschrift fur Vitaminforschung.
Z. Wirls, Zuckerind	Zeitschrift der Wirtschaftsgruppe Zuckerindustrie (Vereins der
a n	deutschen Zucker-Industrie).
Z. wiss. Biol.	Zeitschrift für wissenschaftliche Biologie.
Z. wiss. Mikrosk	Zeitschrift für wissensehaftliche Mikroskopie und mikroskopische Technik.
Z. wiss. Phot	Zeitschrift fur wissenschaftliche Photographie, Photophysik und
	Photochemie.
Z. Zuchtung	Zeitschrift fur Zuchtung.
	Zeitschrift fur die Zuckerindustrie der Čeehoslovakischen Republik.
Zavod. Lab	Zavodskaja Laboratorija.
Zavod. Lab Zellstoff u. Papier	77 11 . M
Zement	Zement.
Zentr. Bakt	Zentralblatt fur Bakteriologic, Parasitenkunde und Infektionskrank-
	hoiten.
Zentr. Min	Zentralblatt fur Mineralogie, Geologie und Palaontologie.
Zentr. Zuckerind.	Zentralblatt fur Zuckermdustrie.

A DICTIONARY OF APPLIED CHEMISTRY.

C—continued.

are methods of saving time in calculating which although well known in some circles are not, it seems, so widely used as they deserve. The object of this article is to give a selection of methods, mainly of technical interest, used for

various purposes.

Weighing by Swings.—A graph may be kept in the balance case on which is plotted the deflection eaused by 1 mg. at different loads. On paper ruled in tenths of an inch with milligrams as ordinates and scale divisions as abscissæ, lines are ruled from 0 through the points indicating the deflections caused by 1 mg. at loads of 100, 50, 10, 1 g. Reference to the appropriate line will enable the final mg. of the weight to be read off without calculation.

Simplified Division of the type 124.712/99.926 may be carried out by adding 0.074 to the denominator and a proportionate number x to the numerator so that the fraction becomes (124.712+x)/100. If $x=0.07\times12/10=0.084$, the quotient is 1.24796; if $x=0.074\times125/100=0.092$, the quotient is 1.24804. Calculated with five-figure logarithms the original division yields 1.2480, with seven-figure logarithms 1.24804. Since the fifth and fourth decimal places have usually no significance in technical work, the

approximation is satisfactory.

Assay Ton.—This is an instance of a general method of avoiding calculations by the choice of a suitable weight of sample for analysis; for instance, in the analysis of ores of gold and silver where the amount of precious metal is expressed in oz. troy per ton of 2,000 lb. avoirdupois. If the weigh-out is 29.1666 g., the number of milligrams of silver or gold found is also the number of oz. troy per ton, since this ton contains 29166.6 oz. troy. Similarly, grains of x per gallon of a water may be estimated by titrating 70 ml. of the sample with a standard solution of which 1 ml.=1 mg. x, since 1 Imperial gallon of water weighs 70,000 grains. In each of the examples given 1 mg. is the same fraction of the amount taken for analysis as is the smaller of the two commercial units of the larger.

CHEMICAL CALCULATIONS. There | required to form a mixture of intermediate concentration c is shown by the diagram, which



indicates that the mixture consists of (c-b) parts of A and (a-c) parts of B. The symbols a and b may indicate normality or other measured property which is proportional to the con-centration. Parts by volume may be used when there is no volume change on mixing. B may be water or a pure solvent of concentration 0.

tration 0.

H. Mager (Chem. Ztg. 1910, 34, 865) has published a collection of mixture-formulæ. The quantities of A and B required (symbols as in the preceding formula) for preparing a stated quantity M of a specified intermediate concentration c are given by the equations x=M(c-b)/(a-b), where x= parts of A, and M-x= parts of B. When a given number N of parts of B, have to be made up to a mixture of concentration. B have to be made up to a mixture of concentration c, by adding x parts of A, x=N(c-b)/(a-c).

In the case that the two solutions A and B contain different solutes and that M parts of a mixture are required containing the solutes in the new proportions a'/b', x=Ma'b/(a'b+b'a). The corresponding problem with three solutions A, B, and C, of concentrations a, b, and crespectively when the specified quantity M of mixture is to contain the solutes in the proportions a':b':c' is solved by the equations

$$x_A = Ma'bc/(a'bc+b'ac+c'ab)$$
.
 $x_B = Mb'ac/(a'bc+b'ac+c'ab)$.
 $x_0 = M - (x_A + x_B)$.

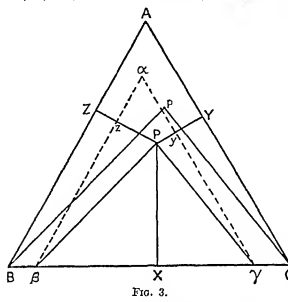
H. G. Nevitt (Chem. Met. Eng. 1932, 39, 673) gives two charts and one table of formulæ for "weighted per cent. calculations," that is, the conversion of weight per cent. into mols. per cent., volumes per cent., etc.

The general formula is $P' = \frac{100}{1 + R([100 - P]/P)}$ Mixtures.—A method of calculating the quantities of two selutions, A (of higher concentration, a) and B (of lower concentration, b) P'=weighted $\frac{P}{N}$ of the lighter component,

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in their study of the relative viscosities of nitrocellulose solutions in mixtures of three solvents used triangular graphs to record the composition of the mixtures, and as shown in Fig. 2 drew lines on the graph joining points corresponding to mixtures possessing the same relative viscosity. The graph shows clearly the effect of alcohol and of benzene in mixtures containing ethyl acetate as the third constituent.

W. Hume Rothery (J. Inst. Metals, No. 2, Graphic Representation by Two Adjacent 1933, 52, 131; C. H. Desch, v. Bibliography) Scales.—The familiar example is a thermometer



(Reproduced by permission of the Institute of Metals.)

gives a graphical method of converting percentages by weight a, b, and c, of three elements, into atomie percentages, the atomic weights being wa, wb, and wc.

The relative numbers of atoms are

$$a/w_a$$
: b/w_b : c/w_c . . . (1) whence

(2) $a/w_a \times w_a : b/w_b \times w_a : c/w_c \times w_a$.

 $a \times 1:b \times w_a/w_b = B:c \times w_a/w_c = C$ (3)

The atomic percentages of the three elements are obtained by dividing 100a, 100B, and 100C, in each case by

a+B+C

In the graphical method, with 60° triangular ruled paper, the percentage composition of the alloy is represented by the point P in the equilateral triangle ABC of height 100 units. The weight percentages of A, B, and C are given by PX=a units, PY=b units, and PZ=c units (cf. Fig. 1). In accordance with equation (3) the points y in PY and z in PZ are then marked so that $Py=PY\times w_a/w_b$, and $Pz=PZ\times w_a/w_c$. Through y and z lines drawn parallel to AC and AB respectively form the equilateral triangle afy. In this triangle the position of the point P

Cochrane and Leeper (J.S.C.I. 1927, 46, 118T) PX, Py, and Pz are in the ratio of equation (3); measurement of these lengths enables the calculation (4) to be carried out. These operations may be avoided by finding a new point p within the triangle ABC geometrically similar to P in the triangle $\alpha\beta\gamma$. This point is found by drawing through B and C lines parallel to βP and γP respectively which meet at p. The atomic composition is given by the position of the point p relative to the triangle ABC.

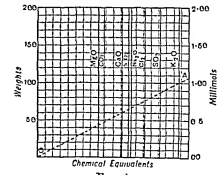
Graphic Representation by Two Adjacent

graduated in Fahrenheit and Centigrade scales. Double scales are easily constructed from an ordinary graph, e.g. from the eurve connecting the vapour tension of water (abscissæ) with the temperature (ordinates). The values of vapour tension for each degree Centigrade are read off and marked on the right hand side of the temperature axis, where they form an unevenly divided scale. Such a scale 22 cm. long for temperatures from 5° to 26°, and pressures from 7 mm. to 25.4 mm., graduated to 0.2°C. and 0.2 mm. is sufficiently accurate for ordinary pur-

Nomograms (v. Bibliography).-In many eases an equation is solved by drawing three (or more) vertical axes graduated with appropriate numerical or logarithmic scales. In use a ruler is placed on the diagram to join observed values on the marginal scales; the imaginary line intersects the third axis at a point on its scalo giving directly the required pertals.) centage, volume, etc. The accuracy obtained is about that of a slide rule for an axis of the same length but may

be increased as required by drawing a portion of the graph on a larger scale.

Useful charts have been designed by H. G. Deming (Ind. Eng. Chem. 1916, 8, 264; v. also



F10. 4.

Bibliography); of these Figs. 4 and 5 have been selected. Fig. 4 is a chart for the conversion of a given weight of one substance to the equivalent weights of a number of others; it solves the equation y=ax for different values of a. corresponds to the atomic composition, since The ordinate scale on the left-hand margin

mixing two liquids (cellulose lacquers) of viscositics 2 and 4 respectively, a straight edge is placed on the chart connecting the points 2 and 4 on the marginal seales, the intersection on the line marked 3 gives the proportion 33.3%

for the 2 poise lacquer.

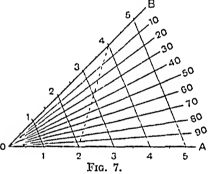
Nomograms in Chemical Analysis.—Zn/ZnO. St. Reiner (Z. anal. Chem. 1926, 69, 377) describes in detail the construction of a nomogram of three vertical axes; readings on the two exterior scales represent the weight of sample and the weight of zine exide respectively, these figures if joined by a black thread or a lino on a celluloid ruler show the percentage figure of zine on the interior axis. The construction of radial charts of the type of Fig. 7 is also described by this author.

S/BaSO₄.—A similar nomogram reading S% on the interior axis was published by O. Liesche (Chem. Fabr. 1928, 1, 161), this graph is drawn advisedly for narrow limits of the weight of sample, and of barium sulphate, enabling 0.1%

to be read between 36% and 44%.

The paper contains an interesting comparison with a chart for the same purpose plotted on

rectangular co-ordinates.



Sb₂S₃ and Sb₂S₅ from S and Sb found.—The diagram replaces calculations in the rubber industry (ibid. 463).

Saponification Values .- A chart in which this number can be read directly is contained in

"Rechenverfahren," 165 (v. Bibliography).
Reaction Velocity.—O. Liesche (ibid. 392, 621, 583) has constructed nomograms in the order given, for the relation between the velocity of a chemical reaction and the temperaturo (chloracetic acid → glycollic acid), the velocity of a monomolecular and of a bimolecular reaction.

Dry Weight % from Wet Weight % .- The chart by the same author (ibid. 377) for this conversion avoids a constantly recurring calculation

in agricultural analysis.

Reduction of Gas Volumes .- Several authors have constructed nomograms for the reduction of observed volumes of gas to N.T.P. Höfsass (J. f. Gasbeleuchtung, 1916, 58, 49) published a diagram which is unfavourably compared by Wa. Ostwald (Angew. Chem. 1919, 32, 359) with his own chart on which the parallel axes for pressure and the reduction factor are divided

with the viscosity 3 poises to be prepared by | by Schwerdt and Loehe (Chem.-Ztg. 1920, 44, 218), who divido the corresponding axes on their chart with a regular scale and as a useful addition give 2 double scales for reducing to 0°C. barometer readings on glass and brass seales respectively. Their temperature scale is curved on account of the correction for vapour tension (v. Barr, infra). Deming (l.c.) like Ostwald employs logarithmic scales, but his chart is too small for practical use. Liesche (l.c. 622) supplies a chart with particulars of the method of construction. F is the sloping axis of correction factors, its scale is projected from the 15° point of the to axis on the left margin, through an evenly divided auxiliary scale. The pressure evenly divided auxiliary scale. The pressure axis on the right margin is graduated downwards from 700 mm. to 800 mm. The t° axis is graduated upwards from 5° to 25°. The upper limit of the F scale F_a reads 0.8437, the lower limit F_c 1.0336. F_a has abscissa $x_{F_a} = 136.1$ mm., Fe has abscissa 127.0 mm. Distance of a point F of the F scale from Fa in mm-

=(1569F-1400)/(0.5566F+1)

Key equation $x_F/(y_F-y_p)=200/(y_t-y_p)$. F = 0.3593p/(273.1+t).

p Scale, $x_p = 200 \text{ mm}$. $y_p = (800 - p) \times 2 \text{ mm}$.

t Scale, $x_t=0$ mm. $y_t=(t-5)\times 10$ mm. F scalo, $x_F = 200/(0.5556F + 1)$.

 $y_F = (1600 - 1548F)/(0.5566F + 1)$.

The length of axes in this chart is about 6.6 cm. A chart for higher temperatures, 200°-1,000°, and pressures 10-200 atmospheres has been constructed (Licsche, ibid. 595).

R. C. Farmer published a "gas calculator," a

nomogram for the correction of volumes of wet and dry gas and also for the logarithm of the weight of 1 c.c. if the gas measured is nitrogen (Baird and Tatlock, London, out of print). Berl and collaborators (v. Bibliography) have drawn a nomogram for correcting the volumes of wet or dry gases. The reduction factor ranges from 0.790 to 1.000 on an axis 21.8 cm. long. Licsche's chart, published by Verlag Chemic (v. Bihliography), avoids the use of a separate axis for wet gases by correcting the volume of a wet gas, measured over water, by means of a vapour tension scale on one side of the temperature axis, which ranging from 5°-26° is 21 em. long. There is also a double seale on the factor axis, the right hand side heing graduated in mg. nitrogen. Both Berl's and Liesche's nomograms, except for their fragile nature, are well adapted for use in the laboratory

The method of constructing a gas reduction nomogram is discussed by G. Barr, who has kindly given his consent to the reproduction of the figure (Fig. 8) and description in his publication (J.S.C.I. 1930, 49, 21T). The following text except for two small verbal alterations intro-duced by Dr. Barr (private communication, 18/2/38) follows the paper cited.

If parallel logarithmic scales are used for B and \hat{f} , the temperature scale is a straight line, divided proportionately to log (1+at), when the gas is dry; when the gas is wet three straight logarithmically. This arrangement is criticised logarithmically divided scales cannot afford an

line, but that for wet gases is curved. The curvature is, however, very slight, and the five points indicated are ample to fix the position and shape of the curve. The suh-division of the t scale into 1/5°C. is most readily performed by

projection.

Fig. 8 is, in reality, two nomograms, of which one scale (B) has been made identical. If the application of the diagram is to be confined to dry or to wet gases only, it may he convenient to add a uniformly divided horizontal scale, as described and figured in J.S.C.I., 1930, 49, 22T, by means of which the multiplication of the

factor may be effected.

Bibliography.—R. K. Hezlet, "Nomography," Woolwich, 1913; H. G. Deming, "A Manual of Chemical Nomography," Nomon Sales Agency, Urhana, Illinois, U.S.A., 1918; J. B. Peddle, "Construction of Graphical Charts," McGraw-Hill Book Co., New York, 1919; S. Brodetsky, "First Course in Nomography," Bell and Sons, London, 1920; W. C. Marshall, "Graphical Methods," McGraw-Hill Book Co., New York, 1921; Swett, "Construc-tion of Alignment Charts," Wiley, New York, 1928; O. Liesehe, "Chemisehe Nomogramme," Verlag Chemie, Berlin, 1929 (cf. articles in Chem. Fabr. 1928, 1929, 1930, 1931); E. Berl, "Tasehenbuch für die anorganisch-ehemische Grossindustrie, 21cr. Teil, Nomogramme," Leibe Grossindustric, 2ter Teil, Nomogramme," Julius Springer, Berlin, 1930; A. Alison, Ind. Chem. 1930, 6, 402 (details of the construction of a nomogram on concentration by evaporation); M. Pirani, "Graphische Darstellung in Wissenschaft und Technik," 2nd edition by I. Rungo, De Gruyter, Berlin, 1931; O. Liesche, "Rechenverfahren mit Anwendungen auf die analytische Chemic," F. Enke, Stuttgart, 1932 (vol. 30 of "Die Chemische Analyse," by B. M. Margosches, continued by W. Böttger); N. R. Corke, "Logarithms Numerical and Graphical," Gee & Co., London, 1934; Van Voorhis, "How to make Alignment Charts," McGraw-Hill Book Co., New York, 1937; C. H. Desch, "Metallo-graphy," Longmans, Green & Co., London, 1937; W. Grosse, "Graphische Papiere und ihre vielseitige Anwendung," C. Schleicher und Schüll, Düren (no date); F. W. Küster—
A. Thiel, "Logarithmische Rechentafeln für Chemiker," 41st-45th edition, W. de Gruyter,

The above classification is Berlin, 1935. J. N. G.

CHEMICAL WARFARE.

Introduction .- Chemical warfare, also called gas warfare, may be defined as the use in war, for the purpose of injuring an adversary, of any substance within the realm of chemistrywhether gaseous, liquid, or solid-which is in any way harmful to the human or animal organism. Chemical warfare is prohibited by the Geneva Protocol of 1925, and the prohibition includes any lachrymatory, irritant, vesicant, asphyxiant, or poisonous substance, hut excludes the normal use of recognised explosives and of products employed as screening smokes, which latter, under normal conditions of use, are not injurious to human beings.

Chemical warfare substances, which are often called poison gases or simply gas, generally on the nature of the gas, but also on the

The temperature scale for dry gases is a straight | produce their effect on the human hody either when breathed in the form of vapour or finely divided particles, or hy contact with the eyes or skin, whether in the form of vapour or liquid.

Physiological Classification.—Chemical warfare substances may be classified according to the chief effect which they exercise on the human body. They fall into the following three main groups, which depend solely on the part of the

body mainly affected:

(1) Lung irritants, which attack the lungs and hreathing passages. The severity of the effects depends upon the substance used. Some gases, such as chlorine and phosgene, will produce death if hreathed in sufficiently high concentration, and are said to he lethal. On the other hand, substances such as the arsenicals, which are used in the form of clouds of finely divided particles (sometimes called smokes) produce intense irritation of the breathing passages, but the effects, while painful, are usually only temporary, and pass off completely in an hour or two after leaving the contaminated atmosphere. This main group is sometimes divided into two classes, namely lethal gases, to include those which may cause death, such as ehlorine and phosgene, and sensory irritants, to eover those which affect the breathing passages hut are generally not lethal, like the arsenicals.

(2) Eye irritants (lachrymators, or gases). Even in very small amounts these have an immediate and painful effect, causing so profuse a flow of tears that it is generally very difficult, if not impossible, to see. The effects soon wear off and no damage is done to the eyes.

(3) Skin irritants (blistering gases, or vesicants). These substances, of which mustard gas is a typical example, attack the skin, and in the worst eases cause deep and extensive blisters. No pain is felt on contact with the liquid or the vapour, and the effects do not show themselves until after a few hours. Mustard gas also attacks the eyes and lungs, and here again there are no immediate symptoms. This constitutes one of the greatest dangers in dealing with mustard gas. There have been recent references to a new type of gas called a "prickling" gas. Such a substance, the main action of which is on the skin, would

The ahove classification is of a very general nature, since most gases act on more than one part of the hody. In fact, most war gases affect the eyes, whatever action they may have upon other parts of the hody. The effects produced hy any war gas depend on the concentration of the gas and the length of time a person is exposed to it. By concentration is meant the amount of gas in the atmosphere; this is usually expressed as 1 vol. or part of the gas, assumed to he in the form of vapour, in a stated number of volumes or parts of air, or as so many milligrams of the gas in a litre of air. The higher the concentration, the shorter is the exposure required to produce death or injury. In the case of gases which injure the lungs, a man must, however, hreathe a certain minimum quantity of the poison hefore it will do him any real harm, the amount of harm depending not only

LETHAL AND INCAPACITATING CONCENTRATIONS FOR AN UNPROTECTED MAN.

	centration wh	tration (i.e. con- ich will incapaci- rotected men for		entration for ed men for f:	Concentration which can be breathed in definitely with-	
	2 minutes.	10 minutes.	2 minutes.	10 minutes.	out damage.	
Diphenyl chloroarsine Diphenylamine chloro-	1:7·5 million	1:75 million	1:8,000	1:40,000	1:1,000 million	
arsine	1:7.5 "	1:75 ,,	1:8,000	1:40,000	1:1,000 ,,	
Diphenyl cyanoarsine	1:17 ,, 1:5	1:170 ,, 1:10	1:16,000	1:80,000	1:2,500 ,, 1:100	
Ethyl iodoacetate Chlorine	1:10,000	1:40,000	1:2.000	(1) $1:10.000$	1:175,000	
Phosgene	1:70,000	(2)	1:10,000	1:50,000	1:1 million	
Chloropicrin	1:100,000	1:200,000	1:4,000	1:20,000	1:1 .,	
Chloroacetophenone .	1:5 million	1:100 million 1:25	(1)		1:250 ,,- 1:250	
Bromobenzyl cyanide	1:10 ,,	1:25 ,,	(1)	(1)	1:200 ,,	

(1) In practice a lethal concentration can hardly he obtained in the field with the lachrymators ethyl iedoacetate, chloroacetophenone, and bromobenzyl cyanide.

(2) A concentration of phosgene which is just insufficient to cause immediate coughing or lachrymation can usually be tolerated, since the initial irritation tends to subside. Such an atmosphere is extremely dangerous, as continued inhalation may end fatally.

aspects of chemical warfare agents is outside hursting charge in the head of the sbell just the scope of the present article, but the bibliography (p. 17) contains a list of the more important publications on the subject. For most purposes the "Manual of Treatment of Gas Casualties," issued hy the War Office and obtainable from H.M. Stationery Office, will be found adequate.

Methods of Use.—Chemical warfare, in the now accepted sense, started with the liheration of chlorine from cylinders hy the Germans in 1915. Subsequently other noxious substances were used in shell, trench-mortar hombs, and grenades, and as experience developed, the shell—by which is meant the artillery shell became the chief weapon for the use of chemical warfare agents. The use of gas in artillery shell was practically confined to shell of calibre of 6 in. and under, the most effective calibres being between 4 in. and 6 in. The artillery shell suffers from the defect that the ratio of weight of gas contained in it to the total weight of the shell is low unless thin walled shells made of steel of high tensile strength are used. The quantities of gas contained in typical war shells were approximately as follows in the case of phosgene and mustard gas:

Calibre.	Weight of gas.	% Ratio of weight of gas to total weight of shell.
3 in. (75 mm. nnd	lb.	
18 pdr.)	1.3	6
4.5 in. (105 mm.)	4.25	10
6 in. (150 mm.) .	11.0	11

In the normal gas shell, an air space of about 10% was allowed, to permit of the expansion

sufficient to break it open and distribute the contents effectively. A percussion fuso was employed. Towards the end of the war the Germans introduced the use of a mustard gas shell with a big explosive charge to produce very fine atomisation of the contents, thus dissipating the gas in a highly toxic cloud hut reducing the amount of liquid scattered on the ground and thus decreasing the persistence. For solid arsenicals like diphenylchloroarsine and diphenyleyanoarsine the Germans employed a shell in which the toxic material contained in a bottle was embedded in high explosive. This shell (called hlue cross) was practically a double purpose shell, as it had considerable explosive value as well as liberating a toxic cloud, or smoke, of finely divided particles of the arsenical.

Trench mortar bombs of 3 in. and 4 in. calibre were also used with effect, but they suffered from the disadvantage of short range. For the gas homhardment of an important area the most effective weapon was the Livens projector, a crude mortar with no great accuracy and limited range, hut which had the advantage that large numbers, running into thousands, could be fired simultaneously. The bomh or drum contained a large amount of gas (about 30 lh.) and the ratio of gas content to total weight was high (about 46%). The gas generally used was phosgene. A very heavy concentration could be put down at once over an area and the concentration was so high that troops taken by surprise were killed or disabled hefore they could put on their gas masks.

At the end of the war a new development was being initiated for the liberation of poisonous or irritant smokes by the volatilisation by means of heat of solid arsenicals in stationary appliances generally known as generators. An important object in using such smokes was to of the contents with heat. There was a small | penetrate the gas mask and cause the wearer to

remove it and so exposs bimself to the action of lethal gases used at the same time. The degree of penetration obtainable depends on the size of the particles in the smoke; the amaller the particle the greater is the penetration, but since the actual quantity of material passing through the mask is what matters from the point of view of the aubsequent physiological effect on the wearer, the result is that for each type of mask there is a particular aize or range of eizes giving the best practical results. This represents the condition which gives the maximum amount of toxic material actually reaching the

wearer. The use of gas in aircraft bombs was subsequent to the Great War a fully authenticated case being the employment of mustard gas in air bombs by the Italians in the Abyssinish campaign in 1935-36. Air bombs can have quite thin walls and the percentage ratio of weight of gas to total weight may be 50 and over, thus showing a very high efficiency as compared with the shell. The post war period has also seen developments for the use of bounds by spraying from aircraft (also employed with mustard gae in the Abyssinian campaign) or in other ways involving discharge under pressure or into heated pipes, such as engine

exhausts. Main Characteristics -The table (pp 12-15) gives the main characteristics of all the chemical warfaro agents which have been used. The term "trench-mortar bombs" refers to all short-range weapons, of which the Livens projector was a particularly effective type. The tarm "grenade" refers to bombs for throwing by hand. The symbols used in the table have the following significance .

Lach. = Lachrymatory

Leth. = Lethal
L.I. = Lung irritant

= Vesicant

= Not important T.M. = Trench mortar

While many substances were used by themselves for offensive purposes, others such as prussic acid were employed only in admixture with other materials in order to obtain their maximum efficiency. Other mixtures were used when it was desired to produce more then one effect at the same time; for example, a mixture of trichloromethyl chloroformate, phosgene, and diphenylchloroarsine was used by the Germana to get a moderately persistent lethal effect in conjunction with the highly irritant but nonlethal action of the arsenical; a further intention was to bring about the penetration of the gas maska of the period, which were comparatively poor in their protection against an arsenical toxic smoke, thus causing the wearer to remove breathing passages and so expose himself to the breatums produced by the constituents, toxic action of the other two constituents, toxic action of the other two constituents, action (CH₂CO CH₃) by gaseous or action regarded as of doubtful utility. The man mixtures are set out below. Where the communities are set out below. Where the compositions were approximately constant the procession of benzyl clothoids (q. 1) to benzyl cyanide percentage of each constituent by neight is (C₂H₂CH₂CN) by treatment with additing cyanida (NaCN) with subsequent brommation.

Bromoscetone (80%) and chloroscetone (20%).

Chlorine (50%) and phosgene (50%). Chlorine (70%) and chloropierin (30%).

Chloropicran (65%) and hydrogen aulphide (35%). Chloropierin (80%) and atannic chloride

Chloropienn (75%) and phosgene (25%).

Cyanogen bromide with bromoacetone or bromomethyl ethyl ketone.

BB Dichiorodiethyl sulphide (80%) and either ehlorobenzene or carbon tetrachloride (20%). Dimethylsulphate (75%) and methyl chloro-sulphonate (25%).

Diphenyleyanoarsine (50%) and ethylcarbazole (50%)

Ethyldichloroarsine (80%) and dichloromethyl ether (20%). Ethyldichloroarsine, ethyldihromoarsine, and

dichloromethyl ether, sometimes with methyldsbromoarsme. Ethyl iodoacetate (75%) and alcohol (25%). Hydrocyanic acid (55%), chloroform (25%),

and arsense trichloride (20%). Hydrocyanic acid (50%), areenio trichloride (30%), stannic chloride (15%), and chloro-

form (5%)
Phosgane (50%) and arsenic trichloride (50%).
Phosgane (60%) and stannic chloride (40%).
Trichloromathyl chloroformate (65%) end

chloropicrin (35%). Trichloromethyl chloroformate, phosgene, and

diphenylchlorosreine. Xylyl bromide and bromoketone or benzyl bromide.

In the table on p. 16 are shown a few important chemical warfare agents which have not been used in actual warfare.

Outline of Methods of Manufacture.

 Acrolem: by distillation of glycerol (CH₂(OH)CH(OH)CH₂-OH) with potassium hydrogen sulphate (KHSO₄) or magnesium sulphate (MgSO₄)

2. Allyl southiocyanate : from natural sources, and by distillation of allyl bromids

(CH, CH CH, Br)

or sodide with alcoholic potassium thiocyanate

(KCNS) or aliver thiocyanate (AgCNS). 3. Arsenic trichloride: by the action of hydrogen ebloride on arsenious oxide (As,O1).

4 Benzyl bromide · hy the hromination of

toluene (C₂H₂·CH₃).

5 Benzyl chloride: by the chlorination of toluene (C₂H₃·CH₃).

6 Benzyl sodide: by interaction of benzyl chlorade (q v.) in alcoholic solution, with sodium

or potaseium iodide (Nal or KI). 7. Bromme: from metallic bromides by the the mask because of the irritant action on his action of chlorine, or of sulphuric seid and an oxidising agent.

10. Bromomethyl ethyl ketone: as for hromoacetone, using methyl ethyl ketone

(CH₃·CO·C₂H₅)

instead of acetone.

11. Carhon tetrachloride: by direct chlorinatien of carhon disulphide (CS,) in the presence of a catalyst.

12. Chlorinc: hy the electrolysis of hrine

(sodium chloride (NaCl) solution).

13. Chloroacctone: by the chlorination of acctone (CH3·CO·CH3).

14. Chloroacetophenone: hy the chlorination of acetic acid (CH₃·COOH) and its subseconversion to ehloroacetylchloride (CH₂Cl·COCl) by treatment with thionyl chloride (SOCl₂). The chloroacetylchloride is then condensed with benzene (C₆H₆) hy the Friedel and Craft reaction in the presence of aluminium chloride to give the desired product.

15. Chlorohenzene: hy the chlorination of

benzene (C₆H₆).

16. Chloroform: from acetone (CH₃·CO·CH₃)

hy the action of hleaching powder.

17. Chloropicrin: hy the action of hleaching powder or chlorine on pierie acid

$(C_6H_2(NO_2)_3OH).$

18. Chlorosulphonie acid: hy the saturation of olcum (SO₃) with hydrogen chloride and subsequent distillation.

19. Chlorovinyldichloroarsine: by the interaction of acetylene (C2H2) and arsenie trichloride (q.v.) in the presence of aluminium chloride (Al₂Cl₆) with subsequent fractional distillation under reduced pressure and treatment to convert the higher homologues, namely dichlorodivinylchloroarsine trichlorotriand vinylarsinc into the product desired.

20. Cyanogen bromide: from the interaction of potassium cyanide (KCN) in aqueous solu-

tlon and bromine at low temperature.
21. Cyanogen chloride: hy chlorination of hydrocyanic acid (HCN) in aqueous solution. 22. Dichlorodiethylsulphide: three mcthods:

(a) By the conversion of ethyleno (C2H4) obtained hy dehydrating alcohol vapour (C2H5OH) over a phosphoric acid or other suitable catalyst into chloro-hydrin (CH₂(OH)CH₂Cl) hy hypochlorous acid; the chlorohydrin is treated with sodium sulphide (Na2S) to give thiodiglycol [(C₂H₄·OH)₂S] which, on treatment with hydrochloric acid or thionyl chloride (SOCl2) yields dichlorodicthyl sulphide.

(b) By the interaction of ethyleno and sulphur dichloride (SCI2) with or without a

solvent.

(c) By the interaction of ethyleno and sulphur monochloride (S2Cl2) with or without scparation of the liberated sulphur.

23. Dichloromethyl chloroformato: hy the gulated chlorination of methyl formate regulated

(H.COOCH₃).

24. Dichloromethyl ether: hy the action of phosphorus trichloride (PCl_s) on trioxymethylene [(CH2O)3] in the presence of zinc chloride (ZnCl.).

25. Dimethyl sulphate: hy the interaction of methyl alcohol (CH3.OH) and chlorosulphonic acid (q.v.).

26. Diphenylamine arsenious chlorido: hy the interaction of diphenylamine [(C6H4)2NH]

and arsenic trichloride (q.v.).

27. Diphenylchloroarsine: three methods:

- (a) By the production of triphenylarsine [(C6H5)3As] hy the interaction of chlorobenzene (C₆H₅Cl) and arsenic trichloride (q.v.) in the presence of sodium and its subsequent treatment with arsenie trichloride.
- (b) By the diazotisation of aniline $(C_6H_5 \cdot NH_2)$ and subsequent treatment with sodium arsenite (Na₂HAsO₃) to give phenylarsonic acid (C₆H₅·AsO₃H₂). The acid is reduced by sulphur dioxide (SO2) to phenylarsinic acid (C₆H₅·AsO₂H̃)₂, which on treatment with a further amount of diazotised anilino gives diphenylarsonio acid [CeH5·AsO2H]. This acid on reduction to diphenyl arsenoxide [(Ph2As)2O] and treatment with hydrochlorie acid yields product desired [(C6H5)2AsCI].

(c) By the reduction of phenylarsonie acid obtained as in (b) partly to phenyl-arsenious oxide (C₆H₅·AsO) and partly to phenyldichloroarsine (C6H5AsCl2), so that the resulting mixture when heated in an autoclave will he converted

into the product desired.

28. Diphenylcyanoarsine: hy the treatment of diphenylchloroarsine (q.v.) with sodium eyanide (NaCN) or anhydrous hydrocyanic acid (HCN).

29. Ethylcarhazole: hy the interaction of earhazolo $(C_{12}H_9N)$ and ethyl sulphate

 $[(C_2H_\delta)_2SO_4].$

30. Ethyldibromoarsine: hy the interaction of magnesium ethyl iodide (MgC2H5I) with potassium arsenite (K₂HAsO₃) to give magnesium ethyl arsenite [Mg(C₂H₆)AsO₃] which is then reduced by sulphur dioxide in hydrohromic acid solution.

31. Ethyldichloroarsine: as for methyldichloroarsine, using ethyl sulphate in place of methyl sulphate or by treatment of ethyl chloride (C_2H_8Cl) with sodium arsenite (Na_2HAsO_3) and the reduction of the ethyl arsonic acid (C2H5·AsO3H2) hy sulphur dioxide to ethyl arsenious oxide (C2H5 AsO), which gives on treatment with hydrochloric acid the desired product.

32. Ethyliodoacctate: hy the chlorination of aectic acid (CH3 COOH) and subsequent esterification to give ethyl chloroacetate esterification to give ethyl chloroacetate (CH₂Cl·COOC₂H₅), which is then treated with

sodium iodide (Na I).

33. Hydrocyanie acid: hy the action of acid on metallic cyanides.

34. Hydrogen sulphide: hy the action of acid on metallic sulphides.

35. Iodoacetone: by the action of potassium iodide (KI) on chloroacctone (q.1.).

36. Methyl chloroformate: by the regulated chlorination of methyl formate (H-COOCH,).

LIST OF CHPWICALS WHICH HAVE REEN USED IN WARFARE AND THEIR CHARACTERISTICS

2						CHE	MICAL	WAI	(FARE	·	_					
	2	Remarks	N.I.	Found in small amounts in shell. Not used	by itself.	NI.	NI NI.	NI.	Fairly important.	Powerful fear man.		N.I.		ı	First gas used.	N.I.
CHANGE	•	Meutealising agent	1	I	Alkalıs.	Sulphur in soda solu- tion.	Ditto.	Alkalı or sodum	(a) Sodum hypo beomito and so dum carbonate	(b) Alkahs.	GINAUS.	ı		1	Sodium hyposulphite (thiosulphate).	As for bromoacetone.
LIST OF CHPWICALS WHICH HAVE BEEN USED IN WARRANT AND THEIR CHAMPALEMENTS	si.	Method of use	Grenades and shell.	1	Used in mixtures with hydrocyanic and	Shell, often in admix ture with xylyl bro	Shell Shell	Grenades.	Shell, with or without chloroacetone or cy anogen bromide.		Shell.	Shell and grenades Sometimes with tylyl	bromide.	As diluent for mustard	Cylinders by itself or mixed with phosgette	or chloropicim. Shell with bromoace- tons.
EN USED IN	-	Physiological action	Lach.	Lach Loth.	No special action.	Lach	Lach.		right Light		Lach.	Lach. L. I.		No special	Leth.	Lach.
A WITTCH HAVE BE		Physical characteristics	Mp 88	2 4 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Mp -18	M P -39	Bp 179 Mp. 24 (docom- poses on dis-	ullition)	M p = 54 B p 136 at 725 mm		N P 29	B p 133–134	Вр 146-146	B p. 768	Mp102 Bp346	B p. 119
LIST OF CHPWICKE	ei	Formula.	сн, сн сно	CH, CH CH, NCS	Ascıs	C.H. CH.Br	0,4,5,4,5 0,4,5,4,5	Br ₁	CH, CO CH,Br		C,H, CHBr CN	сн, со снвгсн,	CH1Br CO CH1 CH1	ccı	°i)	CH3.CO CH2CI
	-	Chemical name	Acroloin (Ally l or neral	dehyde). Allyl 1sothiocy anato .	Arsento trichlonda (Ar- remous chloride).	Benryl bromide.	Benzyl chloride Benzyl icclude	Bromine	Bromoacetone		Bromobenzyl cyanide .	("Bromoketone")	Bromomethyl ethyl ke-	Carbon tetrachlonde .	Chloring	Chloroacetone

				C	HEA	TICAL T	VARFARE	•				13
1	N.I.	Cumulative in its action.	Main value is as a sercening smoko.		1	Powerful war gas; .at- tacks all parts of tho body with which it	whether as vapour or liquid; offect very marked on eyes. Vory insidious. No immediate summediate su	develop in 2-8 hours. Also known as Yperito or Yellow Cross.	Not used by itself; present in trichloro- methyl chloroformato.	N.I.	N.I.	Toxic smoke.
i	1	Sodium sulphide.	Alkalis.	Alkalis.	Ditto.	Bleaching powder, sodium hypochlorito or other chloring in or other chloring needs	0		I	Water.	Alkalis.	1
¥	gus. In mixture with hydro-	Shell and T.M. bombs in admixture with stannie chloride, phosgene or trichloromethyl chloroformate; in cylinders with hydrogen sul-	phido or chlorine. Grenndes and T.M. bombs.	Shell, sometimes with bromoacetone or bro-	Shell.	Shell in admixture with earbon tetrachloride or ehlorobenzene as	90170		1	Shell in admixture with ethyldichloro-arsine and methyl-	Shell in admixture with methyl chloro-	Generators.
No special	No special	Leth. Lach.	No special action.	L.I.	L.I.	V. L.I. Leth.			L.I. Lach.	Lach.	Lach. L.I.	r.r.
B.p. 132	B.p. 61.2	M.p. —69 B.p. 112	B.p. 155	M.p. 62 B.p. 61·3	M.p6 B.p. 16.6	M.p. 14.4 B.p. 217-219			B.P. 116	B.p. 105	B.p. 188	M.p. 190-195 B.p. 410
C,H,C	снсіз	CCl ₃ ·NO ₂	SO ₂ CI-OH	CNBr	CNCI	(CH ₂ CI·CH ₂) ₃ S			сісооснсі	(CH ₂ CI) ₂ O	(CH ₃) ₂ SO ₄	C ₀ H ₄ C ₀ H ₄
Chlorobenzene	Chloroform	Chloropicrin	Chlorosulphonic acid .	Cyanogen bromide	Cyanogen chlorido	ββ'-Dichlorodicthyl sulphide (mustard gas).			Dichloromethyl chloro- formate.	Dichloromethyl ether .	DincthylBulphate	Diphenylamino arseni- ous ohloride (phenarsa- zine ohloride).

7. Remarks	Toxic smoke.	Very effective taxle	NI.	NL	N.I.	Also known as B.K.	N.I.	N.I.	N.I. Not used by itself; present in chlorome- thyl chloroformate.
6. Neutralising agent.	1	1	1	Chlonde of lime,	Chloride of hme.	ag.	1	Alkalis or lime.	11
5. Method of use	Shell by itself or maxed	trickloromethylchlo- roformate; can also be used in generators. Shell as admixture with ethylcarbazolo, Can	also be used in gene- rators. Shell in admixture with diphenyleyanearsine	Shell in admixture with dichloromethyl ether	ario etnyidiculoro- ariono- Sbell in admixture with debloromethyl	bromoarsine. Shell, T.M. bombs and grenades, often dilu-	ted with alcohol. Shell and T.M. bombs in relatures with	senic trachloride, with or without stanno chloride. Cylinders with chloro-	piera. T.M. bomba.
Physiological action.	I.I.	ij	L.I. (slight)	Leth.	Leth.	Lach.	Letb.	Leib.	Lach. L.I. Lach.
3. Physical characteristics	N p. 43	M P 35	M.p. 68 B p. 190	B p. 192-194	B.p. 156	B.p. 178-180	M.p14 B p. 26 5	B.p61-8	B p. 58 at 11 mm. Lach. B p. 71.4 L.I.
2 Formula.	(C,He)2Ascl	(C.H.),AscN	C,H,CNEt	C,H,A1Br,	C,H,Ascı,	CH11-COOC,H1	нси	s,	CH, CO CH,1 CICOOCH,
1. Chemical name.	Diphenylchloroarame ,	Diphenyleyanoarsine .	N. Ethylcarbasole	Ethyldibromoarsine .	Ethyldichloroarsine .	Ethyl lodoscetate	Hydrocyanio acid (Prus-	Hydrogen sulphide.	Iodoacetone . Methyl chloroformate .
	2 Th. Stein Physiological Method of use Neutrilining skent.	2	Parish Payeshed Payeshed	C. H. A. A. A. A. A. A. A	C ₆ +k ₁ , A ₄ C N ₁ + A ₂ N ₂ + A ₃ N ₄ + A ₄ + A ₄ N ₄ + A ₄ + A ₄ N ₄ + A ₄ + A ₄ N ₄ + A ₄ + A ₄ + A ₄ + A ₄ N ₄ + A ₄ N ₄ + A ₄ +	C ₄ H ₂ A2C N ₁ P ₂ N ₂ P ₃ N ₃ P ₄ N ₃ N ₄ P ₃ N ₄	Capacity Capacity	C ₄ H ₂ ₄ A2C N ₁ P ₂ ₄ P ₂ N ₂ P ₂ ₄ P ₂ N ₂ P ₂ N	Parmala, Parmala,

				CH	IEMICAI	WARFARE.				15
N.I.	N.I. Used to a very slight extent.	N.I.	N.I.	N.I. Important war gas. Delayed offects.		Largely used for smoko effect.	Largely used for smoke	enect. About as toxio as chloropicrin.	Moro persistent than phosgeno.	N.I.
l	Chlorido of lime.	Sodium hypobromito		Alkalis; hexamethylenetotramino (hexamino) (in	first gas mask).	Water.	Alkalis.	l	l	Alkalis.
T.M. bombs and shell mixed with dimethylsulphate.	Shell in admixture with ethyldichloroarsino and dichloromethyl	Shell.	Shell.	Shell. Shell and T.M. bombs, by itself or mixed with chloropicrin;	in shell mixed with stannic chlorido or arsenic trichlorido; in cylinders mixed	with chlorine. Grenades, by itself and in shell and T.M. bombs with chloropicrin, and in shell with phosgeno and in hydrocyanic acid	mixtures. Grenades, T.M. bombs,	and shell. Shell.	Shell by itself or in admixture with chloronicin or with phos.	gene and diphenyl- chloroarsino. Shell, sometimes with bromoketono and benzyl bromide.
L.I.	L.I.	L.I.	Lach.	ri. Leth.		L.I. (mild)	L.I. (mild)	L.I. Leth.	Lach. Leth. L.I.	Lach.
B.p. 132	B.p. 181	B.p. 105	B.p. 209	B.p. 252 B.p. 8·2		M.p33 B.p. 114	M.p. 15	D.p. 40 B.p. 73	B.p. 128	B.p. 216-220
SO ₂ CI·OCH ₃	CH,ASBr,	CICOOCH ₂ CI	C,Hs.NC.CI,	C,H;Ascl,		SnCI	so³	cscı,	cicoocci	CH ₃ ·C ₆ H ₄ ·CH ₂ Br (mixturo of ortho, mcta, and para).
Methyl chlorosulphonato SO2CI-OCH3	Methyldibromoarsino .	Monochloromethyl	Phenylcarbylamino	chlorate. Phenyldichloroarsino . Phergene (Carbonyl chloride).		Stannia chlorido	Sul-	pnuric annyanuo). Thiophosgene	Trichloromethyl chloro- formato (Diphosgene)	Xylyl bromido

December on London was a factor of the same of the sam

	7. Remarks,	Fowerful lachrymator, much used for anti-	A very powerful war gas; product gene-	ββ' - dehlorodivinyl - chloroarsina (CHCli CH ₂ AsCl) (b p. 230), which is not quite so effective, and ββ'β', trichlorotrivinylarsino	(CHCI:CH) ₃ As (b p. 260°, m.p. 33°), which is still less active.
UARE.	6 Neutralising agent	Alkalıs.			Chloride of lime and alkalis.
CHESTONE I RODUCES OF IMPORTANCE NOT USED IN ACTUAL WARRARD.	5 Method of use	Could be used in gene. Alkalus. In shell, TM kombs Cov cylinders. Cov cylinders. Cov cylinders. Cov cylinders. Cov cylinders. Cov bombs.			Could be used in shell and bembs.
MIURIANCE NO	Physiological action	Lach.	, 12. 		v. Lī
T I KODOCIS OF I	Physical characteristics %	Mp 659 Bp 245	M.p13 B.p. 190		B.p. 132
TOTAL STATE OF THE	Formula	C, M, CO CH, CI	CHCI:CH Asci		CH ₃ AsCl ₃
	1. Chemical name	Chloroacetophenone . C.H.CO CH,CI	Chloro v myl-dichloro- arsine (Lewiste).		Methyldichloroarsine .

37. Methyl chlorosulphonate: by the interaction of methyl alcohol (CH3:OH) and sulphury I chloride (SO2Cl2).

38. Methyldibromoarsine: by the action of hydrobronic acid (HBr) on methylarsenious

oxide (see Methyldichloroarsine).

39. Methyldichloroarsine: by the action of methyl sulphate[(CH₃)₂SO₄] on sodium arsenite (Na₂HAsO₃) giving disodium methylarsenito [Na₂(CH₃)AsO₃] which on reduction by sulphur dioxido (SO₂) to methyl arsenious oxide (CH3As O) and subsequent treatment with hydrogen chloride gives the desired product.

40. Monochloromethyl chloroformate: by the interaction of methyl alcohol (CH₃OH) and formic acid (HCOOH), and the regulated chlorination of the resultant methyl formate

(H·COOCH₃).

41. Phenylearbylaminechloride: by the dineet action of chilorine on phenyl carbylamine on plicnyl isothiocyanate (C₆H₅·NC), or (C,H,NCS).

42. Phenyldichloroarsine: see diphenylchloro-

- 43. Phosgeno: by the combination of chlorine and carbon monovide (CO) in the presence of a catalyst, usually charcoal; either diluted carbon monovide, such as in producer gas, or concentrated carbon monovido may be used.
- 41. Stannic chloride: by the action of chlorine on tin.
- 45. Sulphur trioxide: by the eatalytic oxidation of sulphur dioxido (SO₂) derived from burning sulphur or pyrites.

46. Thiophosgeno: by the regulated chlorina-

tion of earbon disulphido (CS,).

47. Trichloromethyl chloroformato: by complete chlorination of methyl formate

(H·COOCH^a)

or of chloromethyl chloroformate (q v).

48. Xylyl bromide: by the bromination of

boiling xylene [C₆H₄(CH₃)₂].

Literature References. — The literature on chemical warfare is now so large (running into well over 1,000 different publications) that it is only possible to give a limited number of refercuces to the more important textbooks. are appended under the following four licadings:

- A. Textbooks (covering historical, defensive, and offensive aspects).
- B. Defensive.
- C. Medical.
- D. Offensive.

BIBLIOGRAPHY OF THE MORE IMPORTANT LITERATURE ON CHEMICAL WARFARE.

A. Textbool's (covering historical, defensive, and offensive aspects).—A. A. Fries and C. J. West, "Chemical Warfare," McGraw-Hill Book Co. (N.Y.), 1921; G 11. Fonikes, "Gas: The Story of the Special Brigade," Blackwood (London), 1934, R. Hanshan, "Der Chemische Krieg," E. S. Mittler, Berlin, 1927; J. Meyer, "Der Gaskampf und die Chemischen Kampfstoffe," S. Hirzel, Leipzig, 1926; A. Izzo, "Guerra Chimica e Difesa Anti-gas," U. Hocpli, Milan, Vol. 111.-2

Russian), Moscow, 1929; W. Untermark, "Chemische Kampfstoffe und die Industriegiftstoffe," O. Meissner, Hamburg, 1933; S. de Stockelburg, "Alerte any gas," Lib. Pazot & Cic., Lausanne, 1935 (lengthy appraisal of this by E. Baum in Z. gcs. Schiess- u. Sprengstoffw. 1936, 31, 171, 204-208); Augustin M. Prentiss (and George B Fisher), "Chemicals in Warfare" (with chapters on The Protection of the Civil Populations and International Situation), Populations and McGraw-Hill Publishing Co., Ltd., London,

B. Defensite.—War Office, "Defence against Gas," H.M.S.O., 1935; D. P. Bloch, "La Guerre Chimique," Libr. Berger-Levrault (Nancy), 1927; French Ministry of Interior, "Practical Instructions on Passive Defence against Air Attack," Ch. Lavauzelle, Paris, 1932; L. Leroux, "La Guerro Chimique," Edtns. Spes., Paris, 1933; Izard, Cilleuls, and Kermarrec, "La Guerre Acro-Chimique et les Populations Civiles," Ch. Lavauzelle, Paris, 1933; Fessler, Gebelc, and Prandtl, "Gaskampfstoffe und Gasvergiftungen. Wie sohutzen wir uns?" O. Gruelin, 1933; H. Rumpf, Gasschutz, E. S. Mittler & S. Berlin, 1932; N. Quartara, "La guerra chimica e la difesa della popolazione civile," Turin, 1933; H. Schossberger, "Bautechnischer Luftschutz," Berlin, 1934.

Air Raid Precautions Handbooks: No. 1, ersonal Protection against Gas, H.M.S.O. Personal Protection against Gas, H.M.S.O. 1936; No 4, Decontamination of Materials, H.M.S O., 1936; No 5, Structural Precautions against Bombs and Gas, In preparation; No. 6, Air Raid Precautions in Factories and Business Premises, H.M.S.O., 1936; No. 7, Anti-gas Precautions for Merchant Shipping, H.M.S.O., 1935; Air Raid Precautions: Tho Protection of Foodstuffs Against Gas, H.M.S O., 1937.

R. Hanslian, "The Gas Equipment of Foreign States: Service and Civilian Masks of Various States," Gasseh. u. Luftsch. 1935, 5, 40, 65; P. Murphy, "Defence of the Civil Population against Gas," Listener, 1934, Aug., p. 271.
C. Medical—G. B. Vedder, "Medical Aspects

of Chemical Warfare," Medical Aspects of Chemical Warfare," Williams & Wilkins U.S.A., 1925; U.S.A. War Dept, "Medical Aspects of Gas Warfare" (Med. Dept. of U.S. Army), Washington, 1926; Official History of War: Medical Services—Diseases of the War, Vol. II, H.M.S.O., 1923; War Office, Manual of Treatment of Gas Casualties, H.M.S.O., 1920.

Air Raid Precautions Handbooks: No. 2, First Aid for Gas Casualtics, H.M.S.O., 1936; No. 3, Medical Treatment of Gas Casualties, H.M.S.O., 1937.

L. Dantrebande, "Les Gaz Toxiques," Masson & Cic., Paris, 1933; C. Hederer and M. Istin, "L'Arme Chimique et ses Blessures," Bailhère et Fils, Paris, 1935; H. Buscher, "Grün- und Gelbkreuz," R. Himmelheber, 1932; H. Bucscher, "Giftgas! und Wir," R. Himmelheber, 1932; H. Zangger, "Die Gasschutzfrage," Huber (Bern.), 1933; O. Muntsch, "Leitfaden der Pathologie und Therapie der Kampfgaserkrankungen," Leipzig, 1936; A. Lustig, "Fisiopatologia e clinica dei gas da combattimento," 1930; J. Fishmann, "Chemical Warfare" (in Milan, 1931; A. Lustig, G. Revida, and G.

Ferralora, "Effetti dei gas de guerra (ad uso dei designed. The main exception was the French medici)," Milan, 1934; Il Magne and D. Coedier, M 2 mask, which covered the whole face (eye-"Les gaz de combat au point de vue physiologique, medical et militaire," Baillière et Fils,

D. Offensire.—A. Muller-Kiel," Die chemische Waffe im Weltkrieg und jetzt," Verlag Chemie (Berlin), 1933; M. Sarton, "Chimica del Sostanze Aggressive," U. Hoeph (Milan), 1933 (translated into German by 11. Klumb and published by Vieweg & S., 1935); S. I. Skliarienko, "Physical and Physico chemical Principles of Chemical Warfare '(in Russian), Moscow, 1934. E. Reyes Sarz, "Teora general de la guerre quimica," Toledo, 1933 : D S Sommerville, "The Field Artillery and Chemical Warfare," Field Artill J. 1932, 22, 140. J. T. Hackmann, "Trend of Research for New Chemical Warfare Materials," Chem. Weekblad 1934. 31, 366,; H. Stolt-zenberg, "Die Ultragifte," reviewed in Ind Chem 1935, 11, 460, 483; H. Engelhaed, "Theoretical Considerations on Conventrations of War Gases in the Field," Gasseb. u. Luftseb 1934, 4, 174; R. Baggo, "Artillery Fire with Chemical Shell," Rivista di Artigheria e Genio, 1930, Jan, p. 88; H. Herbet, "Volatility of a Series of Organic Compounds," Koll. Chem Besh, 1926, 23, 313,

J. D. P. CHEMICAL WARFARE DEFENCE. Gas Defence,

Historical and General.-The basis of defence against gas is the respirator, or gas mask When the employment of gas in warfare was instinted by the use of chlorine in 1915, protection for British troops was obtained by the use of pads of flannel, cotton wool, or cotton waste placed over the mouth and nose and soaked in a solution of sodium thiosulphate (hypo) or of this com pound with sodium carbonate. Subsequently the gas mask took the form of a helmet soaked in a solution of hypo, sodium earbonate, and glyrerin, covering the whole of the bead and fitted with mica windows. With the threatened use of phosgene, an impregnating solution of sodium phenate was used for the gas helmet (P. helmet), and was subsequently amproved by the addition of hexamethylenetetramine (hexamine) (P H helmet), which also gave pro-tection against hydrocyanic (prussie) acid. The use of tear gas introduced the need for wearing goggles in addition to the helmet, but the combination was not very satisfactory. As gas warfare developed, the employment of chemical neutralising agents for protection against the variety of gases used and the increasing con centrations became impractirable with respirator of the helmes type, and a gas mask, called the box respirator, was produced by the consisted of an air-tight mask or covering for the good. face fitted with eye pieces of mica, celluloid, or

pieces being provided) and consisted of a number of layers of mushin impregnated with various absorbent chemicals. One mixture contained hexamine, nickel sulphate, sodium carbonate, glycern, and water, and another castor od, alcohol, sodium hydroxide, glycerin, and water. Such masks are very inferior to the face-piera

and container type. Modern gas masks are practically all based on the principle of a face-piece connected to a container of nentralising, absorbing, or filtering materials. The differences in the various designs are essentially differences in details introduced with a view to securing greater service efficiency m one direction or another.

Respirator Face pieces .- Fare pieces may be of many kinds. There are full masks covering the eyes, nose and mouth, and half masks which do not cover the eyes. The latter are useless where gases which irritate or injure the eyes are present; therefore the full mask is always preferable, because, furthermore, it is likely to be more gas tight and comfortable. There are also hoods covering the whole head. The main parts of a face piece are :

(1) The body, generally of rubber, with or without a covering of stockinette or other elastic textelo material on the outside, It esn also be made of leather, of rubberised fabrie in single or multiple layers, of eanvas with or without edgings of rubber, leather, etc., to give a gastight joint, or of asbestos. There are also designs in transparent celluloid or cellophane, so that the wearer's face can be

(2) Eye pieces, generally round, but sometimes oval and fitted with discs of celluloid or of aphnterless, laminated, or plain glass, to the mende of which may be attached gelatine discs to absorb moisture and to prevent fogging due to the mosture in the expired air. The fogging may be obviated by using what is called the Tissot principle, by which the inhaled air passes over the eye pieces and prevents con-densation of moisture. Alternatively or in addition, anti-dimming paste may be applied to the glass. A typical anti-dimming paste can be made from Turkey red oil and water glass For mibtary use, it is important that there should be an adequate field of vision and that it should be possible to use optical instruments such as bmoculars and range finders.

(3) Head harness for adjustment to the head, usually consisting of elastic or spiral metal springs covered with fabric. The bands are generally adjustable to suit the wearer's fare. A well fitting and gas tight mask should not British to deal with all types of gases. This feel uncomfortable on the face if the design is

(4) A rulse holder, for an outlet valve or aplinterless glass; to this mask was attached alternatively an attachment in metal for the by a connecting tube the box or container m container or the connecting tube. The outlet which were placed the materials for the purification of the air breathed by the wearer. This Designs in metal are known, but these are type of construction with a face-piece con generally for inlet valves, which may be in the nected directly or by a tube to a container conatitutes the basic principle on which practically outlet valve is one of the vital parts of a all gas masks for war purposes have sure been respirator as a leakage might be fatal to the

in designing types which are leak-proof under all conditions, and at the same time give good speech transmission. In respirators with directly attached containers there is often no outlet or inlet valve, and the wearer breathes in and out through the filtering box. Certain advantages are claimed for this in the way of increased protection, but it is generally more oppressive to

Face pieces are generally made in three sizes to fit all kinds of face, but a well-designed mask of normal size should, by proper adjustment, give complete protection to well over 90% of adult male faces. Types of face-pieces with nose clip and monthpiece, which were largely used in the war but were very uncomfortable, are now

practically obsolete.

Tho design of a respirator face-piece has to be a compromise between a number of conflicting requirements. It must be gas-tight and comfortable to wear. It must not unduly interfere with vision or with speech, whether direct or through telephones or voice pipes, and it must not impede the wearer in the course of his duties. With the best designs of gas mask the loss of service efficiency is less than

Respirator Containers.—The main chemical interest is in the container, sometimes called the box, drum, or canister, which holds the filtering materials. Containers are generally rectangular or round, and in the normal design, the air enters at the bottom and passes through the filtering and absorbing material, which is generally arranged in horizontal layers. In a few designs, the air enters at the sides, passing through the filtering materials to a central tube connecting with the face-piece. It is not necessary to give details of the internal mechanical structure of a container which can be seen from diagrams, except to say that it is no easy matter to arrango the filtering materials so that they will be effective and remain so under the various conditions of rough handling they may receive in use.

The basic constituent in most respirator containers is active carbon or charcoal, the efficiency of which as an all-round adsorbent for gases and vapours has steadily improved since its first utilisation for gas defence in 1915. The charcoal is used in the form of granules containing generally from 15 to 25% of moisture. This moisture enhances the value of the charcoal for the catalytic decomposition of gases such as phosgene, but reduces its adsorptive capacity for organic vapours. So great is the all-round efficiency of active carbon that some military respirators rely on it entirely for gas protection. It is not, however, active against carbon monoxide, which for physical and physiological reasons is not likely to he used as a chemical warfare agent, nor is it very effective against oxidisable gases such as arsine (arseniuretted hydrogen) or prussic acid. A special mixture of copper and manganese oxides known as hopealite can, however, be used for the

wearer. Great ingenuity has been displayed jeium chlorido in front of the hopealite which soon loses its activity in moist air. The protection against arsine and prussic acid can be improved by the use of granules generally of an alkaline nature and consisting, for example, of soda limo in combination with permanganate. Such granules were used in the British respirator, and not only gave good protection against these two gases, but also against acid gases generally. The value of active carbon as a generally protective agent against poisonous gases can always be enhanced by other types of chemical granules designed to improve the neutralising properties of the container against any particular chemical warfare agent employed. The effectiveness of charcoal as a protection against phosgeno can be greatly increased by impregnating it with hexamine. Such chemical granules are to be found in the containers of respirators designed to deal with special toxic gas hazards in industry (see below). The granule is in effect an active chemical neutralising agent for the particular gas which has to be encountered. Granules may he of two distinct types. The actual chemical or mixture of chemicals may be compressed and then broken up into suitably sized picces, or fragments of a porous material, such as pumice, kicselgubr, or brick, may be soaked in the chemical and dried.

> Neither active carbon nor chemical granules afford protection against fine particles such as are present in screening and toxic smokes, and all service respirators incorporate a filtering material of some sort for dealing with such smokes. These filters vary considerably in composition and efficiency. The principal substances used for making filters are cotton wool, merino wool, animal hair, wood and other paper pulps, cellulose sheets or wadding, feathers, and felt. Their filtering efficiency may he increased by depositing upon them fine smoke particles such as the smoke from hurning camphor, naphthalene, or resins or by mixing them with fibrous materials such as asbestos or with finely powdered lamphlack, or euprene (a solid condensation compound of acctylene) or by depositing nodules of certain gums and resins on them from solution. The difficulty of obtaining a compact efficient filter with a low resistance to breathing has been overcome by various ingenious methods of folding and arranging the layers of filtering material, so that a large filtering area can be compressed into a small space. The best filters now available givo a very high standard of protection against even the most penetrative toxic smokes and will remove more than 99% of the toxic

material from the air breathed.

Industrial Respirators.—Industrial gas masks differ fundamentally from service gas masks in their essential desiderata. The service gas mask has to protect against any concentration that may be encountered of any toxic substance capable of use as an agent of chemical offence. It has, therefore, to be polyvalent. Fortunately, many toxic substances such as carbon monoxide and arsine, against which it is difficult to catalytic exidation of carbon monoxide, if protect by a filtering apparatus, cannot, for required, but this necessitates the provision of technical reasons, be employed in the field. Further, there is a limit to the concentrations

xreptional, while from artillery shell, with which similar material impregnated with caustic soda, nost of the gas attacks were delivered in the potassium carbenate, sodium bicarbonate and/or atter part of the war, a conventration as high | zinc acetate. Active carbon may be used in s 1 in 1,000 is unusual. For toxic smokes, in 1,000,000 is a high concentration. Thus, he problem is one of protection against comaratively low conrentrations of many different sorbent for phosgene; zincates and zincammines. tases.

In industry, the real function of the respirator as an emergency device to deal with unusual reumstances such as the closing down of a eaking valve emitting toxic vapour or the rescue f a man from a dangerous area. Therefore, m ndustry, when a respirator has to be used, it nay be in a confined space and the concentraion may be very high indeed, even as much as few per cent of the toxic vapour. On the ther hand, in any particular works or plant, he danger is generally limited to one or two ases at most. It is therefore possible to design he respirator specially for the gases to be incountered, and so provide a much higher pro ection against them than is possible in a sollyvalent respirator. This means that for ndustry a number of different types are required o cope with all its gas hazards.

The service respirator may have to be worn in action for long periods of time; consequently t must be very comfortable to wear and provide good vision and good facilities for ordinary and elephonic speech. An emergency device, as in ndustry, does not need the same standard of comfort for short period wear, or even of vision or speech, though obviously these should all be good On the other hand, the gas tightness of the industrial respirator must, if anything, be superior to that of the service type because of the higher concentrations. The service respirator therefore is a different problem from the industrial, and is not necessarily austable for meeting the hazards of industry.

The face-piece of the industrial gas mask calls for no special comment, as it is very similar to that used for service purposes. The main difference is in the container. Different types indication of the types of neutralising or absorb-

industrial gas masks.

(1) Organic vapours, such as benzole, acetone, gases

(2) Acid yases, such as hydrocyanic send, example, of sodium thiosulphate pentahydrate, nitric acid, formic acid, free halogens, nitrous are interposed. It is stated that 1% of carbon fumes, phosgene, and sulphur dioxide. Among monoxide will on oxidation give off enough heat the adsorbents used are (a) aods home or other to ruse the temperature of the air adiabatically

hat may be met. A concentration of 1% is alkaline granules, (b) diatomite, brick, pumice or addition to these adsorbents.

Other adsorbents used are based on potassium metasilicate; bexamine, which is a good ad-Soda lime granules impregnated with permanganate will give enhanced protection against hydrocyanic acid, as will also hopcalite, the carbon monoxide neutraliser referred to later. Sometimes a special container is designed to deal with hydrocyanic acid and other weak scids. Diatomite impregnated with finely-divided mercury oxide or certain complex zinc salts is used for this purpose.

(3) Ammonia, the main adsorbents being (a) ealts of copper, such as copper sulphate impreg. nated on pumice (often known as kupramile). (b) salts of zine and cobalt, (c) salts of copper and mckel, and (d) silica gel.

(4) Hydrocyanic acid and carbon dioxide .-- A good adsorbent is baryta granules,

(5) Sulphuretted hydrogen .- A good adsorbent is a mixture of baryta granules and active carbon.

(6) Netrous fumes - Alkaline permanganate granules and hopealite are very useful

(7) Sulphuretted hydrogen and ammonia -Adsorbents used are (a) copper chloride on a solid support which can be active carbon or activated mineral salt, (b) silica gel, (c) mixture of baryta granules, charcoal and copper sulphate granules

(8) Carbon disulphide, for which lead and manganese perovides can be used. (9) Carbon monoxide, against which the gas mask of the fighting services affords no protection. This gas is present in producer gas, blast-furnace gas, coal gas, water gas, afterdamp in mines, etc. As carbon monoxide in any quantity is very often associated with a lack of oxygen, rehance is generally placed on a selfcontained oxygen spparatus when men have to go into such an atmosphere. Of recent years, of container are generally produced to deal with however, much attention has been devoted to different industrial gases or types of gases or for the development of a filter mask for carbon combinations of types. The following is an unnoxide. This has involved the production of a material for the catalytic oxidation of carbon ing materials used for various purposes in monoxide at ordinary temperature. The prodact in most general use is called hopealite, which is a mixture of manganese dioxide and copper earbon tetrachlorude, carbon dissulphute, form-aldehyde, and toluene. The usual adsorbent is other moming are has to be dried, for example, is active carbon or charcoal. Against most by a layer of calcium chloride or since act. organic vapours it is exrellent, but it is poor Behind the hopcalite there has to be a layer of against methyl bromide and methyl chloride; alkaline material, such as soda lime granules, to provided it is not too dry, it functions fairly remove the carbon dioxide formed in the oxidawell against acid gases such as phosgene. It tion of the monoride. The bopcalite will act is asserted by one authority that active carbon indefinitely, so the life of the container depends will stop effectively all gases and vapours with a on the capacity of the drier. Much beat is molecular weight greater than 50. Sinca gel generated in the onidation and the container can in some cases replace active carben, will get very hot if there is much over 1% of but its best value is against water soluble monovide in the inspired sir, while the air will get too hot to breathe unless cooling layers, for

equal to half the volume of the carbon monoxido so that where there is much earbon monoxide the oxygen content in the purified air is likely to be low.

Hopcalite has remarkable properties as an oxidising agent, and will also deal efficiently with ammonia, hydrogen cyanide, hydrogen sulphide, arsine, nitrous fumes, phosphine, and ethylene. Its main defect is its sensitiveness to moisture, which at present is its limiting feature.

Another adsorbent for earbon monoxide, known as hoolamite, is based on the oxidising properties of iodino pentoxide and concentrated sulphuric acid. The mixture is generally used impregnated on pumice with a subsequent layer of active carbon or alkali granules to absorb the

iodine liberated in the oxidation.

As carbon monoxide is tasteless and odourless and produces no immediate symptoms to reveal its presence, the question of a detector to indicate when a container is becoming exhausted is of first-class importance. With most other gases there is ample warning because of the odour, taste, or irritating effect produced from tho first traces of gas that penetrate the respirator; nevertheless, in somo designs visiblo detectors depending on a colour change are fitted. The problem of detecting carbon monoxide has been attacked from many angles, and the following methods for indicating when a container should be changed have been used :

(a) Mechanical timer. This is operated by the breathing and depends merely on the number of inspirations the wearer has made through the container, whether there is any carbon monoxide there or not. It will, as a rule, show the container to be exhausted long before it

(b) Hoolamite detector.—In this detector, when the carbon monoxide passes through the oxidising layer of hopcalito and impinges on the hoolamite, it liberates iodino which gives a warning by its irritating action on the nose and throat that the container is almost ex-Alternatively, it has been hausted. gested that the iedine can be made to liberate chlorine which, combining with carbon monoxide, will give phospene which is readily detectable by the wearer.

(c) Calcium carbide.—When water vapour penetrates the hopealite layer-which it will do when the drier is saturated—it will in contact with the earbide liberate acctylene, which gives

a warning odour.

(d) Colour change.—This is obtained by the effect of humidity on various salts, such as dehydrated cobalt chloride, copper sulphate or platinocyanides. These are placed between the drying material and the hopealite and will change colour when the drier is letting water vapour pass, thus indicating that the hopcalite will soon cease to act. Colour changes are not very reliable and are not easy to observe in dimly-lighted places.

Carbon monoxido respirators have not been used to a great extent in this country as there is always the danger, especially in mines after

about 100°C. The oxidation uses up oxygen | with an inadequate percentage of oxygen in the air. They could, of course, get ample warning of an oxygen deficiency by the use of a test lamp. In many cases no danger from oxygen shortage will exist, and then the carbon monoxide gas mask can be used to advantage, as it will impose less strain on the wearer if he has to do work than will the heavier self-contained apparatus. Apart from mines, there are other cases, for example, near blast-furnaces, where a shortage of oxygen will be accompanied by a high concentration of carbon monoxide, but the latter will immediately make itself noticeable by tho rapid heating of the container and thus give a warning.

To stop industrial smokes, dusts and fumes, the filtering materials used in service respirators

can be employed.

Designs have been produced for all-purpose respirators intended for use primarily by firemen who may have to encounter in fire fighting a largo variety of gases including earbon monoxide and smoke. Such a container is made up from a combination of the adsorbents roquired to protect against the various hazards, and a typical container of this sort on the market consists of the following layers from the bottom to the top; (a) active carbon for organic vapours; (b) cotton wool for smokes, fogs, and dust; (c) caustic soda impregnated on pumice for acid gas; (d) cotton wool for smokes; (e) fused calcium chloride as a drying layer;

(f) hopealite to exidise carbon monoxide; (g) silica gel or carbon impregnated with copper sulphate to stop ammonia. Various other combinations are of course possible. A general adsorbent for gases other than carbon monoxide could be based on sodium carbonate, zine acetate, magnesium peroxide, and powdered

charcoal.

Protection for Animals.—Respirators have also to be provided for animals such as horses, dogs and pigeons in their baskets. These generally take the form of a bag or covering of thick material impregnated with a chemical solution such as was used in the early gas helmets, but they are not very satisfactory in practice.

Collective Protection.—The problem of providing protection for closed rooms and compartments in which people can work without gas masks is solved by the use of purification units constructed on the same general principle as the container of the respirator, through which air is drawn into the room or compartment by means of a fan, preferably from an area likely to be free from gas such as the top of a high building or a chimnoy stack. The methods of air regeneration used, for example, in submarines can also be employed, but are in general less convenient and more expensive. For the protection of dug-outs and rooms in the fighting area, curtains of thick cloth impregnated with a chemical solution or a heavy mineral or a vegetable oil such as cylinder or linseed oils, are used at the entrances, two curtains with an air lock between being generally provided for greater security.

Protection against Mustard Gas .- The introan explosion, that the rescuers may lose their duction of mustard gas with its action on the lives in the event of their encountering a region skin presented a new problem in protection which has not jet been properly solved. Gar- | what simpler and cheaper respirator colled tha ments of oilskin and other impersious materials, civilian duty respirator will suffice. For tha if of air-tight design, give a fair measure of protection, but cen only he worn for comparatively short periods in view of the stilling effect on the wearer, especially in hot weather. The use of clothing impregnated with a chloricating ogent such es chloramine or alhed products which will destroy mustard gas has also con siderable limitations, while protective oint-ments of veseline or of zine oxide and fat mixtures, with or without a chemical to neutralise the mustard gas, ore elso of doubtful utility under service conditions, though they may be of some value in certain circumstances.

Decontamination -Another problem in gas defence is the decontamination or cleaning of oreas which have been contaminated with a persistent gas, especially if it is of the mustard gas type, against which e gas mask is not a complete protection because of its general action on tha skin Generally speaking, water, end carth are the only materials available in adequata quantity if extensive decontamination has to be done, and they can be used to wash away the mustard gas liquid or cover up the infected erea Where available, bleaching powder diluted with about 3 times its voluma of an mort material such as sand or earth is tha most effective aubstance to use The neutralis mos agents mentioned in column 6 of the table on pp 12-15 on chemical warfare agents can also he used for dealing with the gaves against which they are effective decontempation of textiles and general clothing a number of methods can be used depending on the type of garment and the extent of the con tamination, auch as exposure to the weather, washing, solvent extraction, or treatment with atcam or hot air.

indicate the presence of gas While detecting deal with substances such as chlorino and mustard gas, the devices as a rule are not sufficiently simple, quick, or certain in their action to make their adoption desirable except unifer special conditions and in the hands of trained observers Reliance has in practice to be placed on the sense of smell and the sensory

organs generally Protection of the Crest Population against Gas -One of the most important problems of the present day is the protection of the civil popul lation at large against gas ettecks from the air. These attacks may be made with bombs containing gas, or by spreving liquids of high boding point, like mustard gas, from circraft. Those members of the community who must continue to carry on their duties either during or immediately after a raid and the members of the defensive organisations must be provided with gas masks, in the same way or service personnel, and with e certain emount of protective elething depending on the nature of the work they have to do.

civilian population at large the main defence will be gas protected rooms in their own houses or offices, the rooms heing selected with due regard to the fact that high explosive and incendiary bombs ore certain to be used as well as gas. Public shelters, ventilated with air purified by passage through filtration units, will be required for people cought in the streets, and similar chelters mey be installed by important organisations. The public will have to seek the shelter of their gas protected rooms immediately the oir raid warning is given. As a second line of defence to be used if the gas protection of the room is demaged or if e contaminated aree has to be evecuated, the public will be provided with a simple but effective design of gas mask called the general civilian respurator.

In eddition, arrengements will here to be made for :

(a) a gas detection service to examine the bombed areas, diagnoso the neture of the gas used (if any), and indicate whether decontamination is required

(b) decontamination aquads to render the

bombed areas safe to occupy, (c) a first aid service to deal with casualties, which must include first aid and decontamination centres for the treatment of tha Jess serrous casualties and the decontammetion of the persons and clothing of people who may be contaminated with gas, and

(d) casualty clearing stations and hospitals for the more severe cases.

These services will form part of the general air Deletion of Gas -Another problem of roid precaution services required to deal with chemical interest is the use of detectors to all types of air roid and the detailed organisation all types of air roid and the detailed organisation will be the responsibility of the local authorities. papers, paints or solutions can he provided to The necessary technical details will be found deal with substances such as chlorino and in the series of Air Raid Precautions Handbooks which are issued periodically by the Air Raid Precautions Department of the Home Office and are obtainable from H M Stationery Office.

CHEMICKING t. BLEACHING. CHEMILUMINESCENCE. - Accepting the term "luminescence" to denote emission of light other than that due to temperature effects, chemilumineseence " should denote the direct conversion of the free energy of chemical reaction into light energy. As, however, mere change in the energy lavels of ctoms mey result in change of chemical properties end thus be considered as chemical action, end as possibly all radiation of wave length chorter than the mira red is due to release of this store of intre atomic energy, e logical application of modern views would include all forms of luminescence as chemiluminescent phenomena The usual but un-defined terminology therefore restricts the term to those instances when light appears as a result Those of them, like the police and the fire of the formation of molecules of high energy brigades, who will have to perform their duties levels during the production of new substantes, whatever the conditions, will be equipped with In this sense the inclusion of, for exemple, such service respirators. For the others, a some- phenomena as crystallo- and tribo luminescenco

that the colour is the same for both emissions when they can be observed in a single salt (Weiser, J. Physical Chem. 1918, 22, 439) a similar explanation should hold for triboluminescence, and its inclusion among chemiluminescent phenomena in the abovo modified sense should then depend on the demonstration of chemical differences between crystals capablo of exhibiting triboluminescence and aged erystals no lenger possessing such ability. Again, in addition to the traditional use of the term "phesphorescence," this term is also used to denote the delayed emission of light by a substance after previous illumination, fluorescence being the instantaneous reversal of illumination, and it is difficult to decide whether such phenomena are dependent on chemical action in the above sense. Thus Chakravarti and Dhar (Z. anorg. Chem. 1925, 142, 299) were unable to confirm the suspected relationship between fluorescence and oxidation of certain organic dyes, but a chemiluminescent interpretation is not easily disproved in other eases. The luminescence of slightly impure zine sulphide on bombardment with a particles is another border-line instance.

The liheration of part of the energy of interaction between molecules or atoms in the restricted sense may be the result of one of two meelianisms (Wegscheider, Rec. trav. chim. 1923, 42, 585): (a) the energised atoms or molecules emit radiation directly, or (b) the excess energy may be transferred to other molecules which can then emit their own characteristic radiation. Although these conceptions have been experimentally realised in some instances, the mechanism of the majority of known instances of chemiluminescence is quite undetermined, our knowledge being limited to the visual observation of light. The following examples are therefore arbitrarily classified according to the physical state of the

lumineseing system.

The luminescent exidation of phosphorus vapour is among the best known of the phenomena under review (Chariton and Waita, Z. Physik, 1926, 39, 54), although still not completely understood (Downey, J.C.S. 1924, 125, 347; Emeléus and Purcell, ibid. 1928, 628). F. Schacherl (Gazzetta, 1932, 62, 610) finds that phosphorus exychloride exerts a positive catalytic effect and postulates the formation of the exidence of the exidence of chemistry of the emission of light. Aerial exidation of many organic vapours, e.g. ether, earbon disulphide, etc., produces "cold" luminous flames, while the light of the "nonluminous "Bunsen flame is a true instance of chemiluminescence. Oxidation of a 2% mixture of heptane in air producing visible radiation (H. Beatty and G. Edgar, J. Amer. Chem. Soc. 1934, 56, 102), the reaction between hydrogen and chloroform (H. Fromherz and H. Schneller, Z. physikal. Chem. 1933, 20, 158), polymerisation of acetaldehyde in presence of active hydrogen (H. Urey and G. Lavin, J. Amer. Chem. Soc. 1929, 51, 3286), reaction of sodium vapour with chlorine (Lialikow and

is questionable; light emitted during crystallisation of, say, sodium chloride may be ascribed to recombination of ions and as it seems significant that the colour is the same for both emissions when they can be observed in a single salt reactions in the gascous phase.

Terenin, Naturwiss. 1926, 13, 83; Z. Physik, 1926, 40, 107), the "afterglow" of active nitrogen (Bonhoeffer, Z. Elektrochem. 1926, 32, 536), etc., are all further instances of luminescent reactions in the gascous phase.

Some inorganic reactions in solution are known to be chemiluminescent, e.g. many oxidations, the catalytic decomposition of hydrogen peroxide by silver, platinum, etc., and the neutralisation of some oxides by conc. sulphuric acid. The great majority of known eases, however, are furnished by the oxidation of organic materials. Many aldehydcs (Radziszewski, Ber. 1877, 10, 321; 1883, 16, 597), sulphur compounds (Dclépine, Compt. rcnd. 1912, 154, 1171), Grignard compounds (Lifschitz, Helv. Chim. Acta, 1918, 1, 472; Dufford, Calvert and Nightingale, J. Amer. Chem. Soc. 1923, 45, 2058; 1925, 47, 95; Evans and Diepenhorst, ibid. 1926, 48, 715) are luminescent on direct oxidation or on reaction with picryl chloride (Wedckind's reaction), that of phenanthryl-9magnesium bromide being particularly intense (W. Bachmann, J. Amer. Chem. Soc. 1934, 56, 1363). Many dyes and phenolic compounds exhibit the phenomenon when oxidised with hydrogen peroxide or ozone in presence of ferrous sulphate (N. Biswas and N. Dhar, Z. anorg. Chem. 1928, 173, 125; 1930, 186, 154; 1931, 199, 400). The emission of light on oxidising organic hydrazides is not uncommon, e.g. benzhydrazide, benzene sulphonhydrazide, and substituted derivatives (A. Witte, Rec. trav. chim. 1935, 54, 471) that emitted by 3-aminophthalhydrazide being spectacular as it is visible in a dilution of 1:108 an intensity exceeded only by the luminescence accompanying the oxidation of NN'-dimethyldiacridylium nitrate (I)

$$\left[\text{Me(NO}_3) \text{N} \underbrace{ \begin{bmatrix} \text{C}_6 \text{H}_4 \\ \text{C}_6 \text{H}_4 \end{bmatrix} \text{C}}_{\text{T}} - \right]_2$$

which is visible in a dilution of 1:1010 (K. Gleu and W. Petsch, Angew. Chem. 1935,

48, 57) (v. Acridine, Vol. I. p. 130r).

Crystallo- and triho-luminescence have already been quoted as extreme examples of chemiluminescence among solids. Others are the reaction between antimony and chlorine (S. Bhatnagar and K. G. Mathur, Z. physikal. Chem. 1930, 9B, 229), and the oxidation of solid sodium which is non-luminous in pure dry oxygen but becomes luminescent in presence of a trace of water vapour, H₂S, dry HNO₃ vapour, or dry AcOH vapour, but not SO₂ or the halogens (R. Bowie, J. Physikal Chem. 1931, 35, 2964).

Interpretation of certain bright lines in stellar of chemiluminescence. Oxidation of a 2% mixture of heptane in air producing visible radiation (H. Beatty and G. Edgar, J. Amer. Chem. Soc. 1934, 56, 102), the reaction between the Schneller, Z. physikal. Chem. 1933, 20, 158), polymerisation of acetaldehyde in presence of active hydrogen (H. Urey and G. Lavin, J. Amer. Chem. Soc. 1929, 51, 3286), reaction of sodium vapour with chlorine (Lialikow and

on two materians, see germ and receptors, in generalization are every positions of 2022, 102, purified and are remarkably specific, in high 303) studying the oxidation of digitant combined the second components are possible could trace no connection known that where the obtained from the same or very closely related appearance or un non appearance of luminescence species; although considerable incooledge of the conditions governing the luminescence has been of efficiency have revealed at least the over collected, the ultimate structural nature uf these components is unknown. Luciferin has many of the properties of a peptone, and functions with its oxidation product oxy-lociferin as a reversible system. The oxida-tion of luciferin is effected by many oxidising agents hat never with luminescence Luciferase possesses the properties of an albumin and appears to behave somewhat as an enzyme in the liberation of light.

It is unfortunate that the intensity of chemiluminescence is in general so small that the quantitative aspect has been somewhat neg lected. Intensity of emission has sometimes been conveniently determined by visual methods of substitution such as were used by C Thomas and R. Dufford (J. Opt Soe Amer 1933, 23, 251) when studying the chemiluminescence of Grignard compounds, but the sensitised photo graphic plate or moving film presents advantages, eg for recording rapid furctuations in intensity (Amberson, J. Gen. Physiol 1922, 4, 517) Methods employing a photoelectric cell enabling very feeble sources to be studied are particularly valuable, A. Potpelow, B. Pyatnirtki, and S. Zhurkow (J. Russ. Phys. Chem. Soc. 1929, 631 Phys Pt) have studied in this way the luminescence of amarina and lophine when exidised with bromins water. The spectral distribution of the emitted light is determined by spectroscopic means, although when the source is too feeble the use of a series of colour screens of graded limits of transmission yields fairly accurate quantitative results (see Weiser, J. Physical Chem. 1918, 22, 439)

Quantitative study of chemiluminescent phenomena has been concerned chiefly with (a) spectral observations, (b) energetics and efficiency (Bull Nat. Res. Council, USA, 1927, 59, 1). Outstanding conclusions are the recognition of a general band and not a line spectral character; the fact, deducable on theoretical grounds (Harvey, J Gen Physiol on theoretical grounds (marrey, o occu anysome 1925, 8, 89), that for the emission of hight there must occur a state richer by 37-71 cals/mol than another condition to which transition occurs spontaneously. This energy limitation is certainly not the only condition that lumines cence shall occur and may in a complex reaction only apply to an individual stage in the process total energy of reaction should be within these as an anthelmintic for the treatment of bookhmits. Thus the chemiuminescent decomposition of the iodide of Millon's base is actually endothermic (Z. Petrikaln, Z. Physik, 1925, of the efficiency of chemiluminescent processes

of two materials, luciferin and luciferase, in generalisations are rarely possible, e.g. Lifschitz remarkable fact that while the efficiency of " pure " chemiluminescent reactions is in general very small (<1%), that of bioluminescent pro-cesses seems uniformly high. Thus, while the hight efficiency of the (incomplete) combustion uf phosphorus vapour in air was calculated to be 00011% (Adams, Physical Rev. 1924, 23, 771), Karrer estimates the luminescence efficiency of the firefly to be as high as 19 4% (J. Franklin Inst. 1918, 185, 775).

Among a number of reviews of general and special aspects of chemiluminescence may be mentioned the following: F. Schacherl (Atti congresso naz. chim. pura appl., 1933, 4, 599; E Rideal, Chem. News, 1929, 139, 36; Bull. Nat Res. Council, U.S.A., 1927, 59, 1; H. Beutler, Angew. Chem. 1932, 45, 249 (thcoretical), J. H. Graham, Amer. J. Pharm. 1935, 107, 245 (chemiluminescence and biophotogenesu).

CHEMISORPTION P. CATALYSIS IN IN-DUSTRIAL CHEMISTEY. Vol. 11, p. 422r.

GHENODEOXYGHOLIC AGID t. Bille CHENOPODIUM, ESSENTIAL OIL OF. American wormseed oil The oil distilled from the flowering and fruiting plants of Chenopodium ambrosicides, var. antheliminticum A. Gray (Chenopodiacem), a native of Central America and the West Indies. The commercial oil is distilled chiefly in Maryland and is known as Baltimore oil Great care is required during distillation as the chief constituent is decomposed by overbeating High pressure steam is visign as two better activities at the besu as possible. Yield 0 6 to 1 0%.

Constituents - The main constituent is ascaridole (q.v) a body closely related to emeole, having a peroxide structure. Ascaridole decomposes with explosive violence on heating to 130°-150°. Other constituents are p symens, terpinens, \$\Delta^{2-809}\text{menthadiene}, and traces of butyric acid,}

safrole, and methyl salicylate

Characters -A colourless or pale yellow Liquid with a disagreeable, penetrating camphoraccous odour and a burning, bitter taste. Sp gr. 0 960-0 980 at 15 5", optical rotation -4° to -8°, n20 1 474-1 479. Soluble in 3 to 10 vols 70% alcohol. Contains not less than 65% ascaridole, deter-mined by the process described in the "British so that it is not mevitably required that the Pharmacopous." It is employed in medicine worm and round worm.

C. T. B. endothermic (Z. Petrikain, Z. Physik, 1925, CHERIMOYA. The fruit of Anona Cheri-32, 569) It would then seem that estimations molia Mill. It is apparently indigenous to certain districts of Ecuador, and is cultivated of the tenthing of the total heat energy in Menco and other tropical regions of South entitles in lightly can only be of direct America. The fruit somewhat resembles the matching of the first the results of the matching of the first the first tenthing the first mooth of the first the f Even here relationships are so ubscure that modified by the presence of various acids.

1914, 62) recorded the following analysis of the fruit: total pulp 84%, total solids 33.8%, protein 1.84%, fat 0.15%, acids (as malic) 0 09%, reducing sugars 15.34%, sucrose 3.07%, fibre 4.29%, ash 0.67%.

A. G. Po. The fruit of Prunus avium L. CHERRY. (or P. Cerasus, var. avium), the sweet cherry, and of P. Cerasus L. (or Cerasus vulgaris Mill.), the sour cherry. The latter are also known as Morello cherries (dark varieties) or amarcles (light varieties). Cherries are utilised as dessert and culiaary fruit or may be canned, dried or enadied. Marasehino cherries used in confectionery are sweet cherries bleached with sulphur dioxide, split and stoned, dyed with Poaceau 3R, and preserved in syrup. The liqueur maraschino is prepared from the sour cherry. Black Forest and Swiss "Kirschwasser" is distilled from fermented sweet cherries.

The composition of the flesh of cherries, which represents 94-95% of the weight of the whole fruit, averages (% fresh weight):

Ash. 12.0 0.5 0.4 European¹ 14.5 0.51 European² 14.2 9.60.730.6American³ 18.7 0.91 0.43 10.4 0.5

Kulisch, Z. angew. Chem. 1894, 7, 148.
 Ollg, Z. Unters. Nahr.-Genussai 1910, 19, 558.
 Shaw, Oregoa Agric. Exp. Sta. Bull. 1898, No. 55.

Windisch and Schmidt (Z. Unters. Nahr .-Genussm. 1909, 17, 584) record the following analyses of the juice of sour and sweet varieties

1.064 16.8 0.43 1.40 9.32 0.96 0.14 0.5 Sweet 1.067 17.6 0.48 0.64 11.10 0.26 0.09 0.5

According to Hartman and Bullis (Oregon Agrie. Exp. Sta. Bull. 1929, No. 247), the ripening of cherries is associated with an increase in sugar and noa-sugar solids and a decrease in acidity, tanain coateat and astriagency, values recorded being:

	Acids (as maile).	Sugars.	Non-sugar solids.	
Date.	0,0	pn.	6/0	%
June 23	0.75	3.68	7.4	3.4
Aug. 1.	0.72	3.96	16.5	6.2

The sugars consist almost entirely of sucrose, glucose and fructose. Traces of inositol are present. The principal acid is malic acid, and there are very contradictory reports as to the prescace of small amounts of succiaic, citric and tartaric acids.

Traetta Mosca et al. (Annali Chiaa, Appl. 1923, 13. 333) have shown that unripe but not ripe fruit contain a peroxydase, invertase and

Thompson (Hawaii Agric. Exp. Sta. Rept. | The emulsin, according to Hofmann (Biochem. Z. 1934, 272, 426), contains a large proportion of β -glucosidase and very little β -galactosidase. The colouring matter of cherry skins has

been characterised by Willstätter and Zollinger (Annalen, 1916, 412, 164) as keracyanin (chloride, C27H31O15Cl) yielding on hydrolysis with hydrochlorie acid, eyanidin, glucose, and rhamnose. It thus appears to be a diglueoside of evanidin, C₁₅H₁₀O₆.

König gives the composition of the aslı as: K₂O 54.8, Na₂O 4.4, CaO 5.8, MgO 5.4, Fe₂O₃ 1.5, Mn₃O₄ 0.8, P₂O₅ 15.6, SO₃ 5.4, SiO₂ 5.0, Cl 1.6%.

The iron content of cherries appears to vary considerably with the type. Minor elements recorded by various analysts include: Al 34.9, Cu 1.6, Zn 1.5 mg. per kg. fresh weight.

The kernels of cherry stones average about a quarter of the weight of the stone and contain H₂O 6·3%, protein 28·0%, other extract 38·7% (Alpers, Z. Unters. Nahr.-Genussm. 1917, 34, 433).

A. G. Po. CHERRY KERNEL OIL is the fatty oil present to the extent of 30-40% in the seedkernels of the cherry, Prunus Cerasus L. Since 1926, attention has been paid in the United States to the recovery of the oil on a commercial scale from the kernels of the sour cherry, which are available as a by-product of the cherry-canning industry. About 34 tons was thus prepared in 1929, but the potential production of oil, assuming all the pips separated at the eanning plants were utilised for this purpose, has been estimated at about 2,000 tons per annum. Refined cherry kernel oil is pale ia colour and of a bland flavour, and the American product is reported to have good keeping properties. It resembles almond and apricot kernel oils in many respects and is similarly employed in the manufacture of cosmetic and pharmaceutical preparations; it is also suitable for uso as a salad or culinary oil. The following figures have been recorded by Jamieson and Gertler (Oil and Fat Ind. 1930, 7, 371) for a commercial specimen of refined American cherry kernel oil: d^{25} 0-9183, saponification value 190.7, iodine value (Hanus) 115.8, unsaponifiable matter 0.5%; the fatty acids consisted of 2.9% of stearic acid, 4.2% of palmitic acid, 49% of olcic and 42% of linolic acid, together with traces of myristic (?) and arachidic acids. Iodine values from 109 to 123 have been recorded by other observers for crude cherry kernel oils from various sources (cf. Alpers, Z. Unters. Nahr.-Genussm. 1917, 34, 433; Spitaleri, Drug and Cosmetic Ind. 1936, 38, 331; Tilgner, Konserven Ind. 1931, 18,

The oil gives orango or reddish colorations in the Bieber and Kreis tests (see Almond Oil). According to Rabak (U.S. Dept. Agric. Bull. 390, 1916; Oil and Soap, 1932, 9, 210) fresh cherry kernels contain a glucoside of the type of amygdalin, about 1% of volatile oil, resembling essential almond oil (q.v.) being recoverable from the press-cake remaining from the expression of the fatty oil (the kernels from enuls in being present at all stages of growth. imported cherries, which had been preserved by

treatment with sulphur diovide and brine, 399; W. A. Tarr, The Origin of Chert and however, did not have the characteristic bitter Flint, Univ. Missouri Studies, 1926, 1, no. 2; recovery of the fatty oil, could be used as a Great Butain, Mem. Gool. Survey, vol. 6, fertiliser, or, if freed from volatile nil (hydro- Refractory Materials: ganister and silica rock, cyanic acid) as an ingredient of eattle-food.

CHERRY-LAUREL, ESSENTIAL OIL OF. Obtained by distillation of the leaves of Prunus Laurocerasus Linn, (Fam. Rosacex), a native of Persia, and cultivated generally as an ornamental shrub. The leaves contain a cyanogrnetic glucoside laurann which interacts with an enzyme yielding benzaldehyde and hydrocyanic acid. The oil separates from the

98% of benzaldehyde. The distillation water is atandardised to contain 01% of hydrocyame acid, and is used in medicine under the name cherry laurel water, principally in eye lotions

Lauro-CHERRY-LAUREL LEAVES, ceran folia (BP), leaves of Prunus Laurocerasus

Linn. CHERT, A compact form of sules essentially the same as fint, differing from this mainly in its geological occurrence. The name fint is neually restricted to the nodules of irregular shape found in the chalk of Cretaceous age, whilst chert more often forms slabby masses, cometimes thick beds, in rocks of all geological periods. It is usually associated with limestones and dolomites, and sometimes a gradual passage can be traced between the two, suggesting that the beds of chert have been formed by the silicification (metasomatic replacement) of himestone Fossils are also replaced by silica, but there are also often present spongespicules and the siliceous tests of radiolaria, in the latter case the ailica is of organic origin; while in others it has no doubt been deposited from solution as colloidal silica, which has later crystallised on a minute scale. The micro crystalline material consists of finely granular quartz and fibrous chalcedony, together with some opalins silies. The fracture of coert is more uneven and aphntery and the broken surface less smooth than in typical flint.

The material varies considerably in sppearance, ranging from white to black in colour, and sometimes with a cavernous textuce. It also veries in chemical composition. Analyses by H. F. Harwood (Gool, Mag. 1923, 60, 177) nf ehert from north Fintshire show SiO, 9445-9783, Al.O. 036-242, Fe.O. 0-028, FeO 0-022, CO, 0-272, C 0-060%. A collection of analyses of chert from American localities is given by W. A. Tarr (Univ. Missouri Studies, CaCO, 32 12

Deposits of chert are worked at Bakewell in Derbyshire, Halkin in Flintshire, and Reeth in Yorkahire, and the crushed material is used in the production of pottery glazes, and for as a food, or for the manufacture of a flour, or grinding and polishing. (H. C. Sargent, Gcol. as a source of a commercial starch. Mag. 1921, 58, 265; 1923, 60, 168; 1929, 66,

almond" taste) The residual seed meal, after Special Reports on the Mineral Resources of 2nd ed , 1920)

L. J. S. CHESSYLITE or AZURITE. Hydrsted basic copper carbonate, 2CuCO, Cu(OH), forming monoclinic crystals of an azure blue colour. Finely crystallised specimens have been found in abundance in an old copper mins at Chessy, near Lyons, in the south of France, and on this account the mineral is often known as chessyhte (Brooke and Miller, 1852); the name datillation water with a yield of 0.05%.

Characters—Sp gr 1000 to 1055, optical characters colour. Sp gr. 38; hardness rotation—0.30 to +0.30. Contama about 33-4. It occurs as an alteration product of chalcopyrite and other sulphide ores of copper in the upper nxidised zones of mineral veins; and it is itself niten altered to malachite, the green carbonate (CuCO, Cu(OH),). Fine crystals are also found at Broken Hill in New South Wales, Taumeb in South West Africa, and at Bisbee in Anzona; at the last named place it occurs, together with malachite, in sufficient abundance to be mined as an ore of copper, It was also formerly mined at Burra-Burra in South Australia. From Arizona come pretty apecimens, with azunte and malachite banded together, which are polished for use in cheap jewellery. Powdered azurste was formerly used as a pigment under the nome " mountain blue," hut this is now replaced by an artificial product

L. J. S. CHESTNUT. Under the general titla

" chestnut " are included : (1) The awcet chestnut, the seed of Castanea sation Mill (or C vesca Gaertn.), the European apecies, of the large American species, C. dentata Borkh, and of two amaller, dwarfgrowing species, C. pumila Mill, and C. alnifolia

Nutt (the American chinquapins). (2) The horse chestnut, the seed of Esculus

Hippocastanum L.

(3) The water or horn chestnut, the horned fruit of Trapa natane (European), of T. bicornis (Chinese), pr of T. bispinosa (Indian).

(4) The Chinese water cheatnut, which is the onion shaped corm of E leocharis fubricea

Schult. The aweet chestnut is eaten roasted, or akinned and minced as a dressing for poultry, etc, or m Italy is prepared as a flour and used for cooking

The horse chestnut, although rich in nutrient material, has a hitter flavour which precludes ste use as a human food. Auld (J.S C.J. 1913, 32, 173) records that the nuts are not markedly 1926, 1, no. 2); one, from Joplan, Missouri, poissnow to animals and may be used in shows SIO, 2946, while a calcarous chert moderate proportions as a cattle food. Segre from Grand Falls, Missouri, shows SIO, 2057, (Chem. 201, 1916, 40, 212) prepares assistated on the cattle from Grand Falls, Missouri, shows SIO, 2057, (Chem. 201, 1916, 40, 212) prepares assistated to the cattle from Grand Falls, Missouri, shows SIO, 2057, (Chem. 201, 1916, 100, 212) prepares assistated to the cattle from Grand Falls. cattle food by boiling the coarsely ground nuts with successive portions of water.

The etacehy kernel of the horn chestnut or Jesuits' nut renders it serviceable directly

The corms of the true water cheatnut are eaten

either raw or cooked by the Chinese and may also ! be used for the preparation of starch.

The average percentage composition of these "chestnuts" (kernels) is:

	H ₃ O	Protein.	Fat.	N.free extract.	Reducing sugars.	Sucrose.	Stareh.	Flbre.	Ash.
Sweet ehestnut (dried) ¹	4·70 14·0	7·44 8·02	2·90 3·54	80·0 69·9	5·36 2·88		50·6 40·0	2·28 2·80	2·68 2·54
Horse ehestnut (dried)3	2.22	12.08	6-26	74.45	2	51	ř.	2.13	2.86
Horn chestnut (fresh) ⁵ Water chestnut (fresh) ⁶	2·6 84·6 79·2	8·9 2·37 1·84	6·3 0·18 0·18	11·39 17·12	4-4 2-25	11.0	38·7 9·14	2·3 0·4 0·7	2·7 0·8 1·0

above, Baker and Hulton record 5.09% pento- in the cotyledons. De Vevey (ibid. 1908, 15, cans in dried horse chestnut and 3.06% in sweet 696) describes a second glucoside, argyrescin, chestnuts. In water chestnut Hemmi has isolated a hemicellulose yielding l-arabinose on hydrolysis.

The bark, and to a small extent other parts, of the horse chestnut tree contains the glucoside resculin (g.v.). Masson (Bull. Sci. Pharmacol. 1918, 25, II 65) records the presence of resculic acid (needles, m.p. 214°-215°) and resculinic acid able analyses give:

In addition to the earbohydrate material cited (yellow needles, m.p. 230°-231°, leworotary) C27H42O12, yielding glucose and argyrin on hydrolysis.

From the sweet chestnut Venezia (Ann. R. Ist. super agrar Portici, 1932, 5, 136-139)

isolated the polyaleohol castagnitol.

	d ¹⁵ .	Sapon. value.	Iodine value.	Relehert- Melssl value.	Polenske value.	Hehner value.	Acetyl value.	Acid value.	Unsapon. matter.	Refractive index.	Analyst.
(a)	0.926	194.5	95.4	1.54		92.5	13.5			1.4747	Chaplet.1
(b)		175.5	99-0	1.01	0.42	92.8		11-67	2.50	1.4710	Heiduschka and Zeileis. ²

Rev. chim. Ind. 1913, 32, 173.

the horse ehestnut, Belozerski and Dubrovskaja (Biochimia, 1936, 1, 665) isolated a globulin hippocastanin, and a nucleo-protein which on hydrolysis yielded amino-acids of similar composition to those from hippocastanin, and nucleic acid from which guanine, adenine, cytosine, thymine and levulic acid were obtained. Colby gives the following analysis of the ash of sweet chestnut kernels: K2O 48.7, Na₂O 1·2, CaO 4·6, MgO 8·1, Fe₂O₃ 0·4, Mn₂O₃ 0·2, P₂O₅ 23·55, SiO₂ 0·18, SO₃ 12·8, C1 0·34%. Small amounts of copper and zine are also present.

A. G. Po. CHESTNUT EXTRACT. The wood of the sweet or Spanish chestnut (Castanca resca),

In an examination of protein constituents of The bark contains more tannin (17%) than the wood, but is not much used. The tree, which grows to a height of 60-80 ft., is ahundant in Italy, the South of France and Corsica, where it forms immense forests, and it is also very common in America.

Freudenberg and Walpuski (Ber. 1921, 54, [B], 1695) isolated the crude tannin and, by hydrolysis with dilute sulphurie acid, obtained a small amount of querectin, traces of gallie acid, sugar (2%, chiefly dextrose), and ellagic acid (15-20%). All these are present in the combined form. No sugar was produced by the action of emulsin, thus indicating that quercetin and ellagic acid are not present as glucosides. Purification of the crude tannin was effected by the action of aspergillus-tannase, then prethough it contains only 3-6% of tannin, is the cipitating with a lead salt and decomposing source of the much valued chestnut extract.

Baker and Hulton, Aaalyst, 1918, 43, 32.
 Leoncial and Maaettl, Staz. sper. agr. Ital. 1911, 44, 66, 113.
 Quoted by Auld (I.c.).
 Baker and Hulton (I.c.).
 Bralmacharl and Chatterjee, Indiaa Med. Gaz. 1927 62, 365.
 Adolph, Philippine J. Sci. 1926, 30, 287.

² Z. Uaters, Nahr.-Geaussm. 1917, 33, 446.

purafied tannin is a yellowish red, strongly acidic 1914, 30, 212) and Chapmon, Perkin and substance, almost monible in ethyl sectate, and Robinson (40.8, 1927, 3015). The sample con-yields only 3%, of ellapic earl when heated for sizted of dail red frequencing a 5 hours with 3% sulphuric ecid. The scidity indicates a molecular weight of about 400 or some multiple thereof. It gives a hine colora some multiple thereon. It gives the tion with iron ealts, but is not precipitated by bromine water and does not contain phlorogincinol. Chestnut tennin thus eppears to be a representative of a new type of tannin and to be closely allied to that of the native (German) oek.

Müller end Zellner (Biochem. Z 1935, 277, 383) describe the precipitation of a philobaphene, empirical formula C22H20O16, from chestnut tannin extract. When oxidised with alkaline peroxide, it yields a product, C1.H2O,2H2O, decomposing at 400°, which resembles but is not identical with ellagic acid Distillation of the phlobophene with zine dust gives anthracene and naphthalene, but no fluorene, whilst by fusion with potassium hydroxide protocatechuic acid, succinic ecid and pyrogallol are produced.

From the leaves of the Castanea tesca, Knrmeier (Collegium, 1927, 273) isolated a tannin from which, hy hydrolysis, 8-9% of chapte and was obtained. The methylated tennin has the composition Ct. Hij O. (OMe), and yields trimethyl citrate on distillation Munz (bbd. yıelda

1929, 499) ohtsined 3 8% of a tannin, a galloylhexose, from the same source

Chestnut is employed almost entirely in the form of extract, the atrength of which varies, but usually contains from 26 to 32% of tannin. The extract is frequently decolourised, and sometimes mixed with quehracho extract and other materials. Chestnut tannin is the tannin which is most largely employed for the dyeing of silk Castanea resca appears to be frequently confused with the horse-chestnat, Asculus Hippocastanum. The tannin derived from this latter is, however, of little or no practical value (cf. Pollak, Collegium, 1913, 291)

A. G. P. end E J. C. CHIAN TURPENTINE. A turpentine derived from Pistacia Terebinthus Lann.

CHIASTOLITE v. ANDALUSITE

CHICA RED and CARAJURA. These are rare pigments prepered by the Indians of Central America from species of Bignonia which are very similar in appearance and may contain as e basis the same colouring metter. According to Crookes (" Dyeing end Cabco Printing," 1874, p. 388) chira is obtained from the leaves of the Bignonia Chica, which the Indians boil with weter, and add some particles of the hark known as "aryane" to the decanted liquid, which causes the precipitation of the colouring matter An interesting résumé of what is known of chica and similar preparations is given by Holmes (Pharm. J. 1901, 12, 595). Lee (J.C.S. 1901, 79, 284) describes the preparation of a pigment from the heartwood of Bignonia Tecoma.

Erdmann (Jahresber 1857, 487) solated a colouring matter from an alcoholic extract of a specimen of chica red said to be derived from the leaves of the B. chica, and assigned to it the

formula CaHaOa.

The colouring matters present in carajura have been studied by Perkin (Proc. Chem. Soc. peculiar camphoraceous odour, wes insoluble in water, alcohol and dilute alkalis, and contained approximately 4% of colouring matter es calcium or magnesium lake, 23% of peaty matter, per-haps "aryane," a wax, and a resin in small relative amount. The mineral matter es esh (38%) consisted mainly of lime, magnesia end silica, together with some potassium carbonete.

The powdered carajura, after boiling with benzene to remove wax, and digestion with dilute hydrochloric acid at 95°, was extracted with boiling alcohol. The residue obtained on evaporation was digested with hoiling henzene, by which means carajurin, the main colouring matter, was isolated. Subsequent extraction of the residue with hoiling acctone yielded a second colouring matter, carajurone, as an

indistinctly erystalline scarlet mass.

Carajurin, C15H2O2(OH)(OMe)2, garnet needles, m p. 2059-2079, is sparingly soluble in hot benzene and elcohol, more reedily so in pyridine It dissolves sparingly in aqueons caustic alkalis to orenge red solutions which become brown on keeping, but does not dissolve in boiling dilute equeous ammonia (distinction from carajuretin). With elcoholic lead acetate earajurin gives a solution which slowly deposits pale red needles, whereas alcoholic ferric chloride gives a brownish violet solution which becomes brown on the addition of an excess of the reagent. Well defined, crystelline, orange red oxonium salts are ohtained with acids, and from these the hase is easily recovered. Brominetion of carejurin compound, C17H12O5Br C,H,O,Br. which orange apperently both a hydrobromide and a perbromide; on treetment with boiling acetone it is changed into dibromocarajurin, C17H18O Brg, bright red peedles. These bromo derivatives do not melt, but become cerboneceous powders at 300°.

In many respects carajumn resembles the anhydrohydroxyhenzopyrenol bases prepered by Bulow and his collaborators. The coloured anhydrohydroxybenzopyranols heve frequently been found to yield colourless derivatives of hydroxybenzopyranols, and by acetylation of carajurin a colourless triacetyl derivative, mp. 142°, of by drated carajurin is obtained. Further, fusion of carajurin with coustic elkah affords p-hydroxybenzoic acid, whilst by hoiling with concentrated aqueous potassium hydroxide, pacetylanisole is produced.

Demethylation of carajurin with hydriodic acid gives carajuretin hydriodide, C15H11O51, which has the composition end properties of a tetrahydroxyflavyhum iodide. Ita constitution was established by the synthesis of ecutellareinidia chloride:

hydroehloride.

The action of cold pyridine on carajurctin hydriodide leads to the formation of carajuretin, C₁₅H₁₀O₅, scarlet needles which become black at 330° without melting. With acetic anhydride and pyridine in the cold a colourless pentaacetyl derivative, m.p. 156°-157°, of hydrated carajurctin is obtained.

In considering the structure of carajurin there are four possible arrangements differing only in the position of the substituents in the

trihydroxybenzene nucleus:

which proved to be identical with carajuretin Of these (II) and (III) are tautomeric and hydrochloride. Would yield the same hydrochloride. The synthesis of this hydrochloride, however, proved that it was not identical with earajurin hydroehloride. The choice between (I) and (IV) cannot yet be made definitely, but (I) is preferable on account of the colour and stability of carajurin.

Carajurone, m.p. 183°-186° (decomp.), dissolves in aqueous sodium carbonate brownish-red solution, the colour of which is neither so intense nor so persistent on dilution as that obtained from carajurin under similar conditions. The ferrie chloride reaction closely resembles that given by earajurin; alcoholic lead acetate gives a maroon-coloured precipitate. The substance combines with strong acids forming oxonium salts, and there can be little doubt that it is a carajurctin monomethyl ether. The perceptible odour of p-acetylanisole observed on boiling a solution of carajurone in con-centrated aqueous sodium hydroxide indicates that the methoxyl is in position 4'.

Dyeing Properties.—Carajurin and carajurone dye mordanted fabrics in almost identical shades, whereas those given by carajuretin are somewhat yellower. The following comparative results

were obtained on mordanted wool:

	Chromium.	Aluminium.	Tin.	Iron.
Carajurin	Dcep maroon	Dull brownish-red	Bright scarlet	Dcep violet-
Carajurone .	Deep maroon	Dull brownish-red	Dull scarlet	Deep violet-
Carajuretin .	Brown	Dull brownish- orange	Brownish-searlet	Brownish-maroon

A. G. P. and E. J. C.

CHICK PEA. An annual legume, Cicer arictinum, extensively grown in India, where it is known as Bengal grain, and in the Mediterranean countries. The seeds form a valuable food for cattle, or, after removal of the husks, for man. They are boiled and eaten as a vegetable or in some countries are roasted and used as a coffee substitute. A preparation of roasted chiek peas forms an important food substance (Lebleliji) in Bulgaria. In India the young leaves are sometimes fried in oil.

The chick pea plant differs from the unjority of common pea species in possessing numerous glandular hairs which sccrete acid material consisting mainly of citrie, malic and oxalic acids (G. R. Milne, J. Roy. Tech. Coll., 1934, 3,

Average analyses of chick peas and of Lebleliji by Zlataroff and Stoikov (Z. Unters. Nahr.-Genussm. 1913, 26, 242) are:

o,	Protein.	Fat.	N-free extract.	Starch.	I'lhre.	Ash.
----	----------	------	--------------------	---------	---------	------

Chick

10-47 22-6 5-08 56-14 49-33 3-09 2-88 Lebleliji . 6·14 24·8 6·09 58·0 57·99 2·21 2·73 to be:

The composition of the dry matter of stems, leaves, and seeds according to Passerini (Staz. sper. agr. ital. 1891, 21, 20) is: (as percentages):

	Protein.	Fat.		Other carbo- hydrates.	Ash.	Nitro- gen.
Stems	6.35	1.8	35.0	49.0	7.81	1.62
Leaves	14.21	4.1	13.9	58.9	8.83	2.27
Secds	26.20	5.2	1.7	63.6	3.30	4.19

The nitrogen distribution of seeds examined by Zlataroff (Z. Unters. Nahr.-Genussm. 1916, 31, 180) is given as: total nitrogen 3.34, protein nitrogen 2·11, nuclein nitrogen 0·10, ammoniacal nitrogen 0.10, peptone nitrogen 0.007, amido. nitrogen 0.01, amino-nitrogen 0.12. Ivanov (Bul. Appl. Bot. Leningrad, 1933, III, No. 1, 3) observes a very wide range of protein content depending on growth conditions. Extreme values were 12.3% and 31.5%, the lowest occurring in soils from which the specific nodule bacteria were absent. The composition of the globulin of chick peas is shown by Narayana (J. Indian Inst. Sei. 1930, 13, A, 153)

Cystine . Tyrosine		. 0-88 . 4 90
Tryptopha	n	. 041
Argunne		(103 (van Slyke) (121 (durect)
Lysine		76
Histidine		1-0

Small amounts of adenine, betaine, and choline were detected by Zlataroff, who also records the nature of the phosphorus compounds as

	٠.	
Protein-P.O.	0 486	
Lecithin .	0.142	
Soluble organic	0 211	
Inorganic .	0.118	
	_	
Total P.O.	0 998	

The seeds also contain appreciable amounts of

mosite and phytin

The fat present in the seeds has the fellowing sp gr at 15° 0 9396-0 9376, characteristics. solidifying point -19 5°, n 25 1 4744, saporufica tion value 240, the acid number of the crude fat is 0 3 to 0 5, sodine value 110-119, Reichert. Meassl value 4 51, Polenske value 1 1, Hehner value 910, ester velue 2395, unsapomfiable matter 0 45%. The separated fatty needs have m p 25°, todine value 129.

Zlataroff (le) records the presence of 0.3% of a sterol elanutosterol (ocetote, m p 128° Passerini (ic) gives the composition of the ash 13. K,O 246, Na,O 129, CaO 445, MgO 1995, Fe,O, 242, P,O, 3956, SO, 338, SIO, 285, and Ci 0 71%.

A G. Po. CHICLE, Chiele er chiele gum, as it is often termed, is the thickened latex of a Mexican tree, Achras Zapota It is used to a considerable extent in the mannfacture of chewing gum. commercial article is a mixture of water soluble gums resins, and mineral matter. Prochathe and Endemann (Pharm. J. 1879, 9, 1045, 1065) give the following analysis of a clean sample of chicle

Soluble in water	٠		25 00	(gum and mineral salts
Alban			33 75	
Fluavil			22 50	
Hydrocarbons so	lubl	e		
in ether			6 00	
Hydrocarbons in				
in ether .			12 75	•
				E. J. I

CHICORY. Cichorsum Intyous In addieen leaves, blanched by growing in During storage of cheery roots mulin is covering with eoil in the open, are partially converted into mulide, and possibly the young reen leaves, blanched by growing in cellan or regetable-the French endire.

or for the detection of common adulterants, e.g. ground suger beet or artichoke. Wynter Blyth quotes the following for fresh roots :

	Cummy	Gh:	Bitter ex-	4-	ulin and	1
Water	mstter	cose.	tractive.	Fat.	fibre	Ash.
77 0	7.5	11	40	0.6	90	08
Ear	her analy	ses l	y Mayer	(Bied	Zentr.	1885,

Celluloss

828) give:

A	bunu	n-				Bitter	
Vater	oids	Fat	Inulin.	Fibre	Sugar	extract	Ash
72	í-l	02	120	14	56	0.5	14
to			to	to	to	to	to
77			173	18	60	0 15	19
Dur	ıng th	eros	sting p	rocess	there is	conside	rable
lastro	cften	of i	entite s	ind th	in Inst	ted pm	hans

contains increased proportions of reducing sugers together with dextrin and caramel. Petermann found pure roested chicory to con

Water-soluble material (74 2%).

32 123 46

An approximate determination of the composi-tion of mixtures of coffee and chicery may be made, according to Leebody, on the basis that the tincterial power of chicory is roughly three times that of coffee.

The characteristic odour of rossting chicory has been exemined by Reichstein and Beitter (Ber. 1930, 63, [B], 816), who record that the velatile matter produced contains acetaldebyde, aretone, diacetyl, By diketopentane, furfuraldehyde, 5-by droxymethyl furfuraldehyde, maltol, furan, methyl and furfuryl alcohols, acette, pyromete and palamite ueds, together with traces of phenols and a neutral oil. These substances may be regarded as derived from carbohydrate materials Tho large proportions of furfuraldehyde and hydroxy. methyl furfuraldehyde which are derived from mulin, together with the absence of volatile natrogen and sulphur compounds, form a marked contrast with the products obtained during the roasting of coffee. Hehner and Skertally have suggested the determination of furfuraldehyde (as is measure of pentosans) for tion to the widespread use of the ground detecting the presence of chicory in coffee. reasted root for admixture with coffee, the The pentosan content of coffee is 5 0-55%, and green plant self forms a serviceable food, and that of chicory 2 5% (Analyst, 1899, 24, 178).

used raw a s salad or bouled and eaten as a into other more easily fermentable products. Wolff and Geslin indicate the presence in the Complete analyses of cheory are seldom roots of an enzyme which effects the transforma-reported, the majority of published data relates ton : mukin -> mukind -> fructose (Compt. rend. to partial analyses, which may serve to deter-1917, 185, 531; 1918, 196, 428). Thus change mine the composition of coffee chicory mixtures may be brought about in a quantitative manner hy heating at 80°-100°, or in an autoclave, with appropriate amounts of mineral acid, (Bobkov, Z. Spiritusind. 1936, 59, 97, 239, 247). In another investigation Wolff (Compt. rend. 1916, 164, 514) records that the roots contain a substance (probably an enzyme) which he terms inulo-coagulase, which ceagulates inulin in the expressed juice. Its action is apparently

specific for inulin.

The bitter principle of chicory is probably a glucoside of fructose and pyrocatechuic aldehyde (Grafe, Biochem. Z. 1915, 68, 1). Zellner and Richling (Monatsh. 1926, 47, 695), in examining the juice of roots, give the formula $C_{18}H_{20}O_6$ to the hitter substance, and also report the presence of a stearin, mannite and tartarie acid. Wolff also described (1899) an optically inactive sugar which he named synanthrose. Grafo (Beitr. Biol. Pflanz. 1935, 23, 336) states that a glucoside occurs in chicory in which inulin functions as the carbobydrate.

Betaine and chobine occur in small amounts in chicory and are not affected by the roasting process. Adulteration of chicory with sugar beet (which contains approximately seven times the quantity of betaine) is thus easily detected (Vondrak, Zentr. Zuckerind Czechoslov. 1929, 53, 366). A microscopical method for this purpose is based on the occurrence of crystals of calcium exalate in certain cells of sugar heet examined after removal of colouring matter by repeated washing with sodium hypochlorite selution (Collin, Ann. Falsif. 1916, 9, 271).

The presence of chicory in coffee may be

The presence of chicory in coffee may be ascertained by determination of the cupric-reducing power of the extract. Roasted coffee contains 1.9-2.6% and roasted chicory 25-27%

of reducing sugars.

The sp.gr. of extracts also offers a means of detecting adulteration in many cases as the following values for the sp.gr. of infusions of various materials with ten times their weight of water indicate:

Spent tan						1.00214
Acorns						1.00730
Peas						1.00730
Mocha coffee					-	1.00800
Ceylon coffee	•			•		1.00870
Java coffce				•	-	1.00870
Costa Rica coffe						1.00900
Native Ceylon c	off	ee	•			1.00900
Brown malt.	•		•			1.01090
Black malt	•	•	•			1.02120
Dandelien root		•	•			1.02190
Red hect	•	•	•			1.02210
Yorkshire cbico	гу					1.01910
Foreign chicory		•	•	•	-	1.02260
Guernsey chieor	У		•			1.02326
Maizo	•	•		•	•	1.02530
Bread raspings		•	•	•	•	1.02630

(quoted by Wynter Blyth, "Foods," p. 360).

The percentage composition of the ash of chicory roots and leaves is as follows:

K₂O N₂O M₂O C₂O Fe₂O₃ P₂O₅ SO₃ SIO₂ CI indicate the kind of treatment to which the Root 40.4 7.7 6.3 8.7 3.0 14.2 9.0 6.0 3.7 material is subjected. The ribhons, after holling Leaves 60.0 0.7 3.2 14.3 — 9.0 9.0 1.0 1.7 in diluto caustic soda, are exposed to the action

Wynter Blyth ("Foods," pp. 359, 1909) gives as the main differences hetween the ash of coffce and of chicory, the following:

	Coffec ash.	Chicory ash.
Silica and sand	/o 	10.7-35.9
Carhon dioxide	14.9	1.8-3.2
Ferric oxide	0.44 - 0.98	3·1-5·3
Chlorine	0.26-1.1	$3 \cdot 3 - 4 \cdot 9$
Phosphorus pentexide	10.0-11.0	5.0-6.0
Soluble ash	3.0	1.74
		A. G. Po.

CHILE SALTPETRE. Sodium nitrate.

CHILLIES v. CAPSICUM.

CHINA CLAY or KAOLIN v. CLAY. CHINA GRASS. Ramie, Rhea. This fibre is obtained from the stem of a plant of the Urticaecæ, which grows to the height of 4-8 ft. and in general character resembles the common nettle, Urtica dioica Linn., hut is devoid of stinging hairs. The plant exists in two varieties, (1) Boehmeria nivea Gaudich., which bears short, silvery hairs on the lower surface of the leaves, and (2) B. nivea var. tenacissima, the leaves of which are green on both sides. The former variety is cultivated chiefly in China and Formosa under the name of "ch'ü-ma" or "tchou-ma," whilst the latter occurs in moro tropical countries, such as the islands of the Malay Archipelago, where it is known as "rhea." The Indian form has been sometimes regarded as B. nivea var. tenacissima, hut Sir Georgo Watt affirmed that it is of the same variety as the Chinese plant. The cultivation of the plant in China is carried on chiefly in the valleys of the provinces of Hunan, Hupch, Kiangsi, and Szechuen.

The fibre resides in the inner hark or hast tissue of the stem. The methods of extraction differ widely from those used in the case of flax and hemp as the encrusting substances are not removed by a retting process but hy chemical means. The commercial "China grass" is propared in China hy hand. The hark is peeled from the stems and the outer hrown skin or pellicle removed from it hy scraping and washing. The process is tedious, the production of a few pounds of the scraped ribhons constituting a day's work. In these ribbons the fibres are firmly embedded in a gummy substance of a pectinous nature which must be removed to obtain the "filasse" or pure fibre ready for spinning. This operation, termed "degumming," is not carried out hefore export, but is effected subsequently by chemical treatment. The process is designed to dissolve and wash out the gummy substances without attacking the ecllulose; at the same time, the fragments of brown pellicle still adhering become detached and are removed by washing. On account of the danger of the fibre becoming injured by the degumming treatment, spinners usually prefer to buy the fibre in the form of ribbons of handcleaned China grass and degum it hy their own process. Many different processes bavo been devised and are usually guarded as trade secrets. The following will, however, serve to indicate the kind of treatment to which the material is subjected. The ribhons, after hoiling

immersed in a both of dilnte acid, the last two processes being repeated until the whole of the gum has been extracted and the white, lustrous filasse ohtamed.

Field .- From two to four or even six cuttings of remie stems can be obtained annually according to the climatic and other conditions of the locality in which the crop is grown. It is granite and from the china-clay rock. It is estimated that from two cuttings the yield of fresh stems per acre amounts to 15-20 tons, giving 1-1 ton of dry scraped ribbons ribbons furnish about 50% of their weight of degummed fibre or filasse, and hence the yield of filasse per acre amounts to approximately

71-10 cwt per annum.

Characters, Uses and Composition.-Ramie fibre is one of the atrongest known. It is extremely durable and is said to be less affected by moisture than any other fibre, but it is some-The fibre has a what lacking in elasticity. brilliant, ailky lustre, can be dyed readily, and is exceptionally long. The filasse consists of isolated ultimate fibres or small groups of fibres The ultimate fibres vary from 75 to 400 mm in length and from 0-02 to 0 075 mm in diameter, the average diameter being about 0-0375 mm. or about twice as great as that of cotton fibre. The fibres taper towards each end, but the ends themselves are rounded. They are thick walled, have a well marked lumen, and bear longstudinal atriations and occasional transverse markings The fibre aubstance consists of nonlignified cellulose

It has been suggested as the result of a study of ramse by X ray methods that the ultimate fibres are composed of chams of long, thus, submicroscopic crystals which be roughly parallel to the fibre axis. These crystals are estimated to be approximately 0.075 long and 0.005 thick (see W. T. Astbury, "Fundamentals of Fibre Structure" (1933), and G. F. Davidson,

J. Text. Inst 1936, 27, P144)

China grass is used extensively in China for the manufacture of fahrics known as "grasscloths" In Europe the fibre is woven into goods of various descriptions, such as lace, curtains, tablecloths, counterpanes, plushes and even clothing materials. It has been used successfully in combination with wool for the production of certain classes of fabrics and has also found an outlet in the manufacture of mantles fer incandescent gas lighting.

CHINA INK or Indian Ink consists of lampblack made into a paste with animal or fish glue

and dried in the form of cakes or sticks of paint. CHINA STONE. A granitie rock having ita felspathic constituent more or less decom-Posed, but not completely kaolinised, and with an absence of coloured minerals, such as biotite and tournaine. As the felipar still return stocks that are applied for the same purpose.

Part of us alkalı, and as some micaccous mineral
is generally present, the stone is fusible; the bearword of Grantic district, Turro, 1878 to disit is largely used in the manufacture of porcehun, and when free felspar is not introduced

of a solution of bleaching powder, and then covered by W. Cookworthy of Plymouth, about 1750. He first noticed it at Tregoning Hill, near Breage, and described it under the local name of growan or moor stone. It is now obtained principally from the neighbourhood of St. Stephens near St. Austell, In this area it occurs as a local modification of the granite, and is sharply marked off from the normal too hard to be worked like the china-clay rock, and has to be blasted and quarried like ordinary granute; in fact it has been used locally as a building stone. The rock requires no preparation, but is sent direct from the quarry te the potteries. (For an account of the occurrence of china-stone in Cornwall, see Mem. Geol. Survey, Bodmin and St. Austell district, 1909; and the Handbook hy J. A. Howe, quoted below.)

The fellowing commercial varieties are recognised:

1. "Hard purple," a hard, white rock with a faint purplish tinge due to the presence of fluorspar.

2. "Soft" or "mild purple," similar to the

last, but softer.
3. " Dry white stone," a soft, white variety.

4. "Buff atone," similar to the last, but iron stained.

The following analyses by W. Pollard and E. G. Radley are given by J. A. Hows, "A Handbook to the Collection of Kaolin, Chinaelay and China stone in the Museum of Practical

chips and chims stude in the state until of Fractical Geology," London, 1914. I. "Hard purple" china stone from Goonvean, near St. Stephens; IL "Buff" china stone from the same locality; III. China stons from Jersey (" Jersey atona

		- V V U U U U I	ry stone
	I.	11.	III.
SiO, .	. 72 28	73-1g	77 06
TIO,	. 0 05	0.06	0.08
A1,0,	<i>14 90</i>	26.13	12 22
Fe,O,	. 050	0.52	0-47
MnO	. 001	0 0 <u>0</u>	0 19
MgO	. 015	0.14	0 19
CaO	. 1-66	0.61	0.16
Na ₂ O .	. 301	2·1g	3 23
K₃Ō .	. 525	441	5 07
Li ₂ O	. 002	0.02	trace?
H,O (over 1		1.81	0-37
H,O (at 105	າ 0.13	0.20	0 39
P,O,	. 0 53	0-45	0 25
F	. 088	023 Fe	5-030
C1	. 009	— cc	100,0
		_	

100-14 99 96

Jersey atone is a decomposed partly knohnised gramte quarmed in Jersey. Gramte pegmatite grainie quarrieu in oriser, commino regimente in France; granite-aplite at Meldon, near Okehampten, Devonshire; porphyrite in Saxony; lipante in Japan; and felaite in China are all

tinguish china stone under the hame petunzile, a word suggested by the Chinese pelun tee. In to forms the only virtually constituent of the the early part of the Bith century a Jesus paste. China stone is sometimes known as missionary named D'Entrecolks, reusing at 'Cornals stone' in consequence of its being Kings-technic sent to Reamunt, in Parts, assiplies largely quarried in Cornwall, whece it was dis- of the raw materials used in China. These were the first specimens of their kind that had ever reached Europe, and they led to the foundation of the manufacture of porcelain at Sèvres. The china stone was described by D'Entrecolles as pe-tun-tse, and the china-clay as kaolin; but it appears that the former name, if not both, must have been erroncously applied. According to the Chinese scholar, Stanislas Julien (Hist. et Fabric. de la Porcel. Chin. Paris, 1856, preface xx.), pe-tun signifies "white paste," while tse is merely a diminutive applied to the material when worked up into small cakes. Hence, petun-tse is really the name of small blocks of white clay or prepared paste, and not of the granitic rock. Nevertheless, custom in this country has justified the use of the term pc-tun-tse as synonymous with china-stone.

L. J. S. CHINA WOOD OIL, nut oil, synonyms for

ČHINAPHENIN. QUINAPHENIN. Quinine carbophenetide,

C20H23ON2·O·CO·NH·C6H4·OEt.

CHINEONAL. Quinine diethylbarbiturate.

Sedative and antipyretic.

(Ger., Porzellanblau.) CHINESE BLUE. Several pigments are sold under this name, of which the following are examples: ultramarine and flake white; cobalt blue and white lead;

and a double cyanide of iron (Prussian blue).
CHINESE GREEN or LOKAO v. LOKAO.
CHINESE RED v. AUSTRIAN CINNABAR.
CHINESE VEGETABLE TALLOW,

Stillingia tallow, is the hard fat which coats the seeds of the Chinese tallow tree, Stillingia schifera Michx. (syn. Stillingia sinensis, Croton sebiferus L., Sapium sebiferum Roxb., Excacaria sebifera F. Müll. Fam. Euphorbiaccw), a tree growing wild, and also largely cultivated, in China, Indo-China and Northern India. Plantations have recently been developed in Flerida and Texas. In Tonkin, the tree is also valued for its leaves, from which a decoction is prepared for dyeing silk. The fruit capsule contains three oval seeds surrounded by a mass of the true vegetable tallow. This "tallow" is to be distinguished from the oil obtained from the seeds (kernels) themselves, known as stillingia oil or locally as "tsé-tiéou" or "ting-yu," which is a liquid drying oil. The tallow may be scraped off the seeds by passing them between fluted rollers, or may be melted off by steaming the seeds in perforated cylinders. The product is pressed into cakes in native hot presses, and sold under the Chinese name of "pi-ićou" or "pi-yu," and is known in Europe as "prima vegetable tallow." According to According to another process, the seeds and tallow are crushed together, giving a mixture of the tallow and stillingia oil, known as "mou-iéou" or, to European candlemakers, as "secunda vegetable tallew"; this product is naturally much softer and has a lower melting-point and much higher

is not suitable for cattle food, as it contains a Greenland. saponin; in China it is used as a fuel.

The properties of the tallow vary considerably according to the source. The refractive index is 1-4546-1-4556 at 40°; the saponification value varies from 197 to 215, and the iodine value from 19 to 32. Considerable quantities of free fatty acids may be present; thus Hilditch and Priestman (J.S.C.I. 1930, 49, 397) found as much as 25% in a specimen from the United States. They analysed this sample, and also two less pure Chinese specimens, and found the fatty acids to consist of 57-69% palmitic, 21-34% oleic, 3-6% myristic, and 1-3% stearic acids. The tallow contained about 25-35% of fully saturated glycerides (largely tripalmitin), and over 60% of mono-oleo-disaturated glycerides (chicfly oleo-dipalmitins).

The solidification temperature appears to vary greatly, being usually about 38°C., but samples, probably impure, have been stated to solidify at 24°C., whilst the American sample examined

by Hilditch set at 48.2°C.

Exports of Chinese vegetable tallow from Hankow to Europe and the U.S.A. amounted to 5,000-6,000 tons in 1923-25, but have greatly diminished since that time, and none has been imported into England for some years prior to 1938. A large proportion of the production is consumed in China for the manufacture of candles; the total exports from Hankow were 17,800 tons in 1917, and 8,200 tons in 1931. In Europe vegetable tallow was chiefly used in the candle and soap industries; its value as a confectionery fat is uncertain (cf. Diedrichs, Z. Unters. Nahr. Genussm. 1914, 27, 132).

J. L. and E. L. CHINESE YELLOW (King's yellow). A mineral pigment owing its colour to the presence

of hydrated ferric oxide.

CHINIOFON. Iodohydroxyquinolinesulphonic acid mixed with sodium bicarbonate; antiseptic for the treatment of amobie dysen-

CHINOFORM. Trade name for quinine

formate.

CHINOSOL or QUINOSOL. Trade name for the potassium salt of 8-hydroxyquinoline-5-sulphonic acid,

C.H.NO.SO.K,H.O,

a pale-yellow crystalline powder with a saffronliko smell and a burning taste; m.p. 175°-177.5°; readily soluble in water, sparingly soluble in alcohol, insoluble in ether; with ferric chlorido gives an intense green colour, yellowish needles with copper salts, and a white precipitate with barium chloride. At a concentration of 1 in 1,000 its solutions possess as great a bactericidal action as mercuric chloride solutions of the same strength. An antiseptic and disinfectant.

CHINOTROPIÑ or QUINOTROPIN.

Trade name for urotropine quinate. CHINOVOSE v. CARBOHYDRATES.

CHINQUAPINS v. CHESTNUT.

CHIOLITE. A double fluoride of aluminium and sodium, 3AIF3.5NaF, closely akin to cryolite When fully developed, the tree yields 25-30 [AIF₃·3NaF], but crystallising in the tetrakg. of fruit a year; the fruit yields about 30% gonal system. Found at the Ilmen Mountains of secunda or 15% of prima tallow. The cake in the southern Urals and with cryolite in

CHIOS TURPENTINE RESIN. character to ordinary turpentine oleo resin. Emmanuel (Pharm. Acta Helv. 1935, 10, 12) isolated from the resin of Pielacia Terebinthus, 1901ated from the result of reacted personaur, terminthe acid, C₁₁H₂₀O₂, m₁, 130°-137°; terminthence acid, C₁₁H₂₀O₄, m₂, 102°; and terminthelma acid, C₁₁H₂₀O₄, m₂, 102°; and terminthelma acid, C₁₁H₂₀O₄, m₂ 128°, These acids have not yet been characterised by the preparation of crystalline derivatives

Chios turpentine is variable in composition, and its characters depend entirely on the relative proportions of essential oil and resin

E. J. P. Chirata, BP. Is the plant CHIRETTA. Swertia Chirata Buch -- Ilam collected when in flower and dried Japanese chiretta is Suertia chinensis Franchet Höhn (Arch. Pharm. 1869, 215) found two bitter constituents in Indian chiretta, viz chiratin and ophehe acid.

CHITENINE, QUITENINE. An oxida tion product of quinine, found in the urine after the administration of quinine Crystallises from dilute alcohol in prisms, m.p. 281°-282°,

[a]_D¹⁷ -122 6°. CHITIN. [a]_D -14 7° (in cone HCl) Isa polysaecharide containing nitrogen which forms part of the skeletal substance of insects and crustacem, it is also an important akeletal element in the fungs. It is not possible to distinguish between animal and vegetable chitin by total nitrogen or by X ray analysis, their chemical identity has been shown by Zechmeister and Toth (Z physiol Chem 1934, 223, 53).

Chitin is extremely resistant to hydrolysis, but on boiling with concentrated hydrochloric and it is converted into I mol of glucosamine (2 ammoglucose) together with I mol of aretic

It is considered by Meyer and Mark (Ber 1928. 61, [B], 1936) to be built up of N aretyl glucosamine units in f glucosidic linkages exactly as in cellulose.

Karrer and Hoffman state that an enzyme from the vineyard enail is able to hydrolyse chitin (Helv Chim Acts, 1929, 12, 616, 986), the end product being acetylelucosamine

By acetolysis of chitin with acetic acid in sulphurio acid, Bergmann et al (Ber 1931, 64, [B], 2436) obtained the octa-acetate of a diascharde chitobiose

Zechmeister and Toth (Ber 1931, 64, fB). 2028, 1932, 65, [B], 161, 1706) obtained in

aoluble chitodextrin.

The Röntgen diagram (Meyer, Helv. Chim.

The | only slowly, but after solution in concentrated name Chios turpentine is, properly, restricted hydrochloric acid, and separation by pouring to the oleo resin from species of Pistacia, all into water, the chitin is easily hydrolysed. though turrentine from some of the larenes is N-Acetylglucosamine is the final product alika often termed Chios turpentine. It is aimilar in from chitin of animal and fungal origin; chitodextrins are intermediate products. Chitinase is not the same as emplain; it has, how. ever, been obtained from the outer part of almonds free from \$ glucosidase (Grassmann, Ber, 1934, 67, [B], 1; Helferich, Z. Physiol

Chem 1933, 221, 253). It has also been obtained from Aspergillus oryzae and is obviously widely distributed. it is destroyed at 70°. The optimal p_{π} is 5.2; Perhaps it is a mixture of two enzymes acting in

auccession.

Chatmase is able to hydrolyse aynthetic glycosides of N-aretylglucosamine, for example, phenyl N.acetylglucosammide. It is quite without action on the non acctvlated compound : chitosan is only hydrolysed as far as the polyglucosamine stage, whereas acetylchitosan is totally hydrolysed by the enzyme. The acetyl group is thus essential for the enzyme to be active; it cannot be replaced by formyl or benzoyl.

CHITOSAMINE is glucosamine (2-amino-

glucose) (v Chutin).

CHIVES, Allium Schanoprasum, L. perennial plant occurring naturally in many parts of Europe and cultivated for the round onion like leaves which are used for flavouring. The percentage composition of the leaves is given as .

N free Water Protein Fat extract Fibre Ash 91.2 26 0.33 3 09 1.48 1 28

Churg and Ripperton (Hawaii Agric, Exp. Stat, Bull 1929, No. 60). The mineral constituents include Ca 0 048, Fe 0 0084, and P 0 057% A. G. Po.

CHLOANTHITE. Native nickel arsenide, NiAs, isomorphous with smaltito (CoAs,) there being no sharp line of demarcation between the two species. Found as cubic crystals and compact masses at Schneeberg in Saxony and Riccheladorf in Hesse, where it was formerly mined as an ore of mekel It occurs in considerable amount with silver ores at Cobalt and South Lorrain in Ontario.

L. J. S. CHLORAL, TRICHLORAGE I ALDE-HYDE, CCI, CHO. Chloral was first obtained by Lachig (Annalen, 1832, 1, 189) by chlorination of absolute alcohol. Its composition was estabhshed by Dumas (Ann. Chim. 1834, [11], 56, 125)

addition a chitotriose and an amorphous water and by Städeler (Annalen, 1847, 81, 101).

Chloral is manufactured by chlorination of absolute alcohol Chlorination is carried out in Acta, 1935, 18, 589) also confirms the structure lead or lead lined vessels provided with a reflux as being of the long chain cellulose type. It is condenser and a chlorine distributor taken to not known whether the glycoside linkage is a the bottom of the vessel, and so arranged that a maximum distribution of chlorine passes through E F. A. the alcohol. The vessels, of from 400 to 1,000 CHITINASE, the enzyme which hydrolyses gallons capacity, are about two thirds filled chitin, was discovered by Karrer and Hoffmann with alcohol, three such vessels being arranged (liely, Chim Acta, 1929, 12, 816) in the digestive in series so that any excess chlorine from the birst juices of Helix It attacks genuine chitin vessel passes into the second and from the

record to the third in order to ensure complete absorption of the gas. The hydrogen chloride evolved during the reaction is absorbed in water. The initial reaction is vigorous and during the first stage the temperature is kept as low as possible by efficient cooling. Chlorine is passed in at a rate which results in a liquid of approximately 24°Bé. at the end of the first day's run. During the next twenty-four hours the temperature is raised gradually, heat being applied if necessary, to about 50°C., and the density of the liquid at the end of this period should be from 35°-40°B6. The reaction is completed on the third day by increasing the temperature to 95° and continuing the chlorination until the density reaches 49°Bé. A sample of the product at this stage distilled with an equal volume of concentrated sulphuric acid should indicate a yield of about 75% of chloral. The erude chloral alcoholato is allowed to cool, when it solidifies. It is then gradually mixed with an equal volumo of sulphurie acid 66°Bé, the mixture being kept cool. The temperature is then gradually raised. Hydrogen chloride is evolved, together with some ethyl chloride. Between 70° and 90° alcohol is recovered, and erude ehloral passes over between 90° and 98°. The erude chloral is purified by redistillation over calcium carbonate, the portion distilling over above 94° being pure chloral.

Other processes which have been suggested include the chlorination of alcohol in the vapour phase (G.P. 133021), the ehlorination of a mixture of acetaldehyde and alcohol (F.P. 612396), and the ehlorination of acetal (Reichert, Bailoy, and Nieuwland, J. Amer. Chem. Soc. 1923, 45, 1552).

Chloral is a colourless, pungent liquid, b.p. 97.7°. When puro it is stablo, but in the presence of traces of impurities such as sulphuric acid it polymerises with production of metachloral, a white amorphous solid. The same product is obtained by the action of aluminium chloride on chloral (G.P. 139392). Metachloral is insoluble in water, alcohol, ether and acids, but soluble in sodium earbonate solution. On distillation at 180°-185° it is reconverted into ehloral (Kolbe, Annalen, 1845, 54, 183). A water soluble polymeride is obtained by treating chloral with pyridine or an amine in the cold and then acidifying. Alcohol and water convert it into chloral alcoholato and ehloral hydrato respectively. Alkalis decomposo it, giving ehloroform and formie acid.

CHLORAL HYDRATE, CCI, CH(OH)2, is by far the most important derivative of chloral. It is prepared by the enutious addition of the requisite amount of water to chloral, overheating of the mixture being avoided. It is purified by crystallisation from benzene, ehloroform or light petroleum. To obtain the hydrate in the form of cubes or plates rather more water is added than is theoretically necessary, the mixture thoroughly shaken until cold and the mass of crystals poured on to poreclain dishes and dried over sulphuric acid in racuo.

Chloral hydrate occurs in colourless crystals m.p. 50°-58°, with a pungent odour and bitter taste, and is readily soluble in water, alcohol,

ehloroform, ether and oils.

Chloral hydrate is very largely employed in medicine as a hypnotic and is official in most pharmacopæins. It is of special value in simple nervous insomnia, delirium tremens, and certain forms of insanity. It is also a powerful deodorising and antiscptic agent. By itself, or in concentrated solution, it may be used as a vesicant. The toxic effects produced by overdoses of chloral hydrate are a fall of temperaturo and slow and enfeebled respiration.

FORMAMIDE, CHLORAL CHLORALAMIDE, C3H4O2NCl3, is prepared by gently heating eliloral and formamide in equimolecular proportions. On cooling the melt sets to a solid mass which is recrystallised from water or 30% alcohol. It forms colourless crystals, m.p. 114°-115°. It is soluble in water (1:20) and very soluble in alcohol, ether and acetone. It is not decomposed by acids, but when warmed with dilute alkalis is decomposed, yielding ehloroform, ammonia, and formie acid.

Chloral formamido is a somewhat slower acting hypnotic than chloral hydrate and is especially useful in the insomnia of cardiae disease, since it has not the depressant action of chloral on the heart. It is also used in combination with potassium bromide as a remedy for sea sickness.

GLUCOCHLORAL, CHLORALOSE, C8H11O6Cl3, obtained by heating chloral and glucose in equal parts on the water bath, forms erystals, m.p. 185°, is a hypnotic and sedative. An isomeric product, parachloralose, produced at the same

time is devoid of hypnotic properties.

BUTYL CHLORAL, TRICHLORBUTYRIC ALDE-HYDE, CH₃ CHCl CCl₂ CHO, is prepared by passing dry chlorine into aldehyde or paraldehyde at about -10° until the aldeligde is saturated. The temperature is then gradually raised to 100°, chlorino being continually passed in until chlorination is complete. The resulting liquid is diluted with water and then distilled in a current of steam, when the hydrate passes over. The hydrate is recrystallised from water and on distillation in a stream of hydrogen ehlorido tho pure chloral is obtained (Pinner, Annalen, 1875, 179, 26). It is a colourless oil with a characteristic odour, b.p. $164^{\circ}-165^{\circ}/750$ mm. sp.gr 1-3956 at $20^{\circ}/4^{\circ}$; fuming nitric acid converts it into triehlorbutyric acid. It readily combines with water forming the hydrate.

BUTYL CHLORAL HYDRATE,

CH3.CHCI.CCI,.CH(OH),,

is prepared by mixing butyl chloral with about one-ninth its weight of water and recrystallising the solid mass so formed from boiling water. It forms white trimetric plates with a pungent but not acrid odour and a nauseous, bitter taste. It melts at about 78° and resolidifies at about 71°. It is soluble in about 40 parts of water, very readily soluble in alcohol, ether and glycerine, less readily in chloroform and olivo oil. Butyl ehloral hydrate resembles chloral hydrate in its action, but is a weaker hypnotic and has a more pronounced depressant action on the heart. It is chiefly employed in combination with eamphor, phenazone or gelsemium as an analgesic in cases of neuralgia and migraine. A. J. E.

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CHLORALAMIDE. formamide, CCI, CH(OH) NH CHO (v. CHLORAL). CHLORAMIN r. ACTIVIS.

CHLORAMINE.T, sodinm ptolnenesulphonchloroamide,

CH. C.H. SO, NCINa 3H,O,

first prepared by Chattaway (J.G.S. 1905, 87, 151), is a white crystalline solid with a slight cdour of chlorine and a hitter taste soluble in about 7 parts of water, and may be obtained from p toluenesulphonyl ehloride, a by product in the manufacture of saccharin, or directly from tolurne by sulphonation and conversion to the sulphonyl chloride, which is then treated with ammonia to form the sulphon-This, by the action of bleaching powder amide or sodium hypochlorite, is converted into the dichloramine, which, in presence of excess of caustic sods, yields chloramine T.

Chloramine T contains about 25% uf "available" chlorine. It was introduced by Dakin during the war as a solution for the arrigation of wounds to replace the much less stable bypo chlorite solution, and is now included in the British Pharmacopous Aqueous solutions of the salt are comparatively stable and may be kept for some time without appreciable decom

position

Technically chloramine. T finds employment in many directions in the textile industry as a hleaching agent and for the development of colour on the fibre, as well as in the manufacture of printing inks and the preparation of soluble starch for finishes, etc. It is also used in the brewing industry for general cleansing and sterilising purposes and as a preservative in the treatment of leather, and in gins and adhesives It does not readily attack aluminium, brass, tin or enamel, and is therefore available for the cleansing and disinfection of vessels used in the preparation and storage of food products.

Dichloramine T, p toluenesolphondichloro-amide, CH₃C₂H₄:SO₄NCl₂, was also pre-pared by Chattaway (le), and is a yellow crystalline powder, m.p. about 78°, having an odonr of chlorine. Unlike chloremine T, it liberates bromine from sodium bromide solution On exposura to air it is gradually decomposed with loss of chlorine. It is sometimes employed in solution in oil as an antiseptic.

A. J. E.

CHLORANIL, Tetracbloro-p-quinone. CHLORAZENE. Syn. for Chloramine-T.

CHLORAZIOE, N.CI v. Azomidz. CHLORENE. (1) A green hydrocarbon of

unknown structure obtained as a by-product in the preparation of fluorocyclene by heating acenaphthene with lead oxide at 350° (Dzieworski and Suknarowski, Ber. 1918, 51, 457). Dilute solutions in benzene are pure green in colour, mora concentrated solutions are purple-red; chlorene solutions undergo photochemical oxidation on exposura to light. The absorption spectrom shows three well-defined bands in the visible region

(2) The term "chlorene" has also been

Syn. for chloral- applied to a yellow gaseous hydrocarbon, C.H. made by passing acetylene through a quartz tube at 750°. Its reactions are similar to those of acetylene, and it is allowly converted at the ordinary temperature to a colourless isomer (Mignonac and Ditz, Compt. rend. 1934, 199, 367).

CHLORETONE. Chlorbutol. Trichloro tert-butyl alcohol, Me CCCl OH. Sedative.

CHLORINE.

Historical.-Chlorine must have been known, according to Mellor (" Inorganic and Theoretical Chemistry," Vol. II, p. 20), to many alchemists from the thirteenth century onwards. It is, however, to Scheele that we owe the first adequate description of the preparation in 1774 of "dephlogistigated marine acid air "ohtained during his researches on the properties of black oxide of manganese,

Lavorsier and Berthollet, a few years later, preferred the name " oxymuratic acid," and the view that it was a rompound persisted until 1809, when Gay Lussac and Thenard suggested it was an element. In 1810, Davy adduced evidence which was regarded as conclusive that the gas was an element, the characteristic colour of which

led him to auggest the name chlorine. The observation of the bleaching properties of the gas end its solutions in water led Berthollet, 1785, to its practical application, and in 1789 to make the still more convenient alkaline bleaching solution. The solution of chlorine in canstic potash became known as Ean de Javelle, whilst the solution in cau, ic soda became Eau de Labarragne The name Eau de Javelle is still widely used, but nowadays refers to a solution of

sodium hypochlorite

The fact that bleachers in Lancashire had used a solution of chloring in milk of lime pre vented Tennant, 1798, from obtaining a valid patent. In 1799, bowever, Tennant patented the use of dry hydrated lime to absorb chloring to make "bleaching powder," and thus was born one of the major heavy chemical industries. Blanufacture of bleaching powder, which was first carned out in Glasgow, spread rapidly as the sikah manufacturers realised its importance, particularly as an outlet for their waste hydrochlorio acid.

The production of chlorine and caustic soda when an electric current passes through a solution of common salt was observed at the beginning of the nineteenth century. Industrial development of this process, however, waited on the supply of cheap mechanically-produced electric energy introduced with the Gramme dynamo electric machine in 1872. Commercial development occurred with increasing intensity from that date.

Occurrence.- Chlorine, apart from volcanio gases, never occurs in nature in the free state (Shepherd, J. Washington Acad. Sci. 1920, 10, 23). In combination with other elements it occurs as:

1. Sodnm chloride in sea water and in rock salt deposits.

2. Potassmm ehloride alone and combined with magnesium chloride or sulphate in salt deposits.

3. The chlorides of calcium, magnesinm,

lead, copper, iron, manganese, mercury and silver.

4. Hydrochloric acid in volcanic gases and springs.

5. Sodium and potassium chlorides in the vegetable and animal kingdom and as hydrochloric acid in the gastric juice.

The chief chlorine supplies are derived from salt deposits formed by the evaporation of inland seas or sections of the sea isolated by change in land levels. Solar salt produced by evaporation of sea water is used to a minor extent as a source of chlorine.

Sodium chloride in the form of rock salt occurs in large deposits in England, Germany, Austria, France, Spain, Russia, Canada and the U.S.A. Deposits occur in various geological formations: in England, for example, the rock salt is found in the Keuper marl, a Triassie formation, whilst the deposits in the States of New York and Michigan and in Ontario occur in Silurian rocks.

Potassium chloride is found associated with other salts in the Dead Sea, and is one of the products of the solar evaporation practised there. The chief sources of notassium chloride are the deposits of Stassfurt, Alsace-Lorraine and Spain, where it is found combined with other salts, as, for example, KCl-MgCl₂,6H₂O,

Laboratory Preparation .- The oldest method of preparing chlorine for laboratory purposes consists in heating manganese dioxide with hydrochlorie acid:

MnO2+4HCl=Cl2+MnCl2+2H2O

The most convenient means of obtaining chlorine in the laboratory is from a cylinder of the liquid. The liquid is not, however, quite pure, containing small quantities of bromine, organic compounds, and dissolved gases, the total of such impurities being less than 10% by weight.

When pure chlorine is required, the method of von Graebe may be used. Analytical quality hydrochlorie acid is allowed to drip on to crystals of potassium permanganate covered with water, a gentle heat being supplied to give a steady evolution of the gas.

Physical Properties.—At. wt. 35-457; atomic no. 17; isotopes 35, 37.

Properties of the Gas.—Density at N.T.P. 3.214 g./litre. This density is about 1.7% greater than required by the gas laws, but very close approximation to the ideal density occurs above 300°C. Above 1,137°C. dissociation into atomic chlorine occurs to some extent (Reinganum, Physikal. Z. 1905, 6, 1514).

The mean value of the coefficient of expansion at atmospheric pressure and ordinary temperature is 0.003836 (Jaquerod and Tourpaian, J. Chim. phys. 1913, 11, 15).

lleat capacity at 15°C., and 1 atm. Cp=

0.115 cal./g. The ratio Cp: Cv=1.355.

The explosive limits of mixtures of hydrogen and chlorine are 91.9% Cl. and 8.1% H. to 14.3% Cl. and 85.7% H., when using an induction coil spark (P. Mathieu, J. Physique, 1917, 7, 166).

Properties of the Liquid.—B.p. -34-6°C. The vapour pressure in atmospheres, the densities of the liquid and the saturated vapour arc (D= g./ml.):

Temp. °C.	P. atm.	D. liquid.	D. vapour.
-34·6°	1.00	1.561	
-20°	1.84	1.524	
-10°	2.61	1.496	
O°	3.65	1.468	0.0128
+10°	4.96	1.438	0.0175
+20°	6.57	1.408	0.0226
+30°	8.60	1.377	0.0300
+50°	14.1	1.310	0.0486
+100°	37.6	1.109	0.136

Critical temperature 144°C. Heat capacity 85.4°C. to -76.1°C.=0.229 cals./g. Surface tension 27 dynes/em. at the boiling-point. Viscosity 0.494 centipoises at -35.4°C.

The expansion coefficient is:

Latent heat of vaporisation at -22°C.= 67.4 g.-cal. at +8°C.=62.7 g.-cal.

F.p. -103°C. with vapour pressure 8.9 mm. Latent heat of fusion 1.626 kg.-cal./g. mol. Solubility.—At total pressure 760 mm. lis the volume of gas dissolved by 1 vol. of water (gas volume reduced to N.T.P.), and q is the weight of gas in g. dissolved in 100 g. of water.

The heat of solution for 1 mol. in 1,000 mols. water is 4.96 kg.-cal. Chlorine is more soluble in 14% aqueous hydrochloric acid and less soluble in salt solutions than in water:

20°C. wt. % HCl 4·13 6·61 8·74 10·40 13·94 20°C. wt. % Cl₂ 0.703 0.722 0.743 0.769 0.823

(Oliveri-Mandalá, Gazzetta, 1920, 50, 94). In saturated NaCl solution:

$$t^{\text{CC}}$$
.
 .
 $14\cdot5^{\circ}$
 $29\cdot0^{\circ}$
 $60\cdot0^{\circ}$

 Wt. % Cl_2 .
 0.0966
 0.0842
 0.0364

Chlorine is also readily soluble in inorganie liquids, such as SiCl4, SnCl4, SO2Cl2, and CrO₂Cl₂, etc.

At 0°C. SiCl, dissolves 13% Cl, by weight. At 0°C. SO₂Cl₂ dissolves 14% Cl₂ by weight. Organic liquids also dissolve chlorine:

Aretic acid et 15°C.:

99 84 90 0 75 0 650 Wt. % Acetic acid

pressure below 96°C, jellow crystals are deposited Moist gas on cooling below 96°C. also deposits the same substance, which is a hydrate of chlorine. The hydrate formed under different conditions of temperature, pressure and relative concentration of the constituents, has the composition Cl,,6H1O (Anwar-Ullah, J.C S 1932, 1176

The hydrolysis of chlorine in water was studied by Jakowkin (1899), who found equilibrium was attained at 25°C, after 48 hours Increase in temperature favours the hydrolysis and decreases the time required to reach equilibrium (Z. physikal Chem 1899, 29, 613).

The photolysis of chlorine in water (Alimand, Cunhife and Maddaon, JCS 1925, 127, 822; earned over by the contaminating gases and 1927, 655) has been investigated, and an lost. Moreover, a limit is set to the cytent of explanation given for the mechanism of the hquefaction by the hydrogen content of the reactions which produce hydrochloric and residual gee, which for safe working must not chloric acids together with oxygen

For more complete physical data see "Comprehensive Treatise of Inorganic Chemistry," Vol II, by J W Mellor, "International Critical Tables", Gmellin'a "Handbuch der anorganischen Chemie System No 6." Landolt

Boernstein, Tabellen

Chemical Properties - Chlorine is a very reactive substence, and even a smell percentage of the gas in air has e characteristic pungent and irritating smell, the danger of breathing chlorine is not that it is a true poison, but that it rapidly ettacks the mucous and lung mem branes, couring congestion

Chlorine combines with most elemente with con siderable evolution of heat and in some cases with ignition, when uncontrolled reaction takes place

Phosphorus ignites spontaneously in the ges, forming phosphorus trichloride Finely divided! arenic, antimony, copper, tin, lead and iron burn in chlorine, forming the respective chlorides, but most of these metals, when in bulk, behave oute differently. The action of chlorine on metals is also profoundly affected by the amount of water present, the amount of reaction decreasing with decreasing water content. The indifference of dry chlorine to ateel enables liquid chlorine to be manufactured and shipped governed by the rules and regulations of railway in steel vessels

Chlorine and hydrogen may be mixed together in the dark without combining, but such mixtures ex plode violently in bright day light or when heated Either alone or in the presence of catalysts, chlorine combines with many inorganic chemical compounds, eg sulphur dioxide, earbon monoxide, phosphorus trichloride, ferrous chloride,

stanneus chloride

forms substitution products and the eliminated hydrogen is simultaneously converted into hadrogen chloride. With benzene and other Wt v_o Cl₂ 106 714 465 381 aromatic organic compounds, eddition or different Hydride — On cooling a solution in substitution products for both are formed water saturated with chloring at atmosphere according to the temperature, and catalyst conditions

LIGHTO CHLORINE.—Liquid chlorine was first obtained to 1805, by Northmore. The pure gas can be condensed to a liquid at 15°C, by a pressure of about 6 atmospheres, or at atmoaphene pressure by reducing its temperature to -35°C. Chlorine gas from electrolytic rells impairties consist of air, drawn in by working the cells under abgbt suction, carbon dioxide, and oxygen from hydroxyl, sulphate and chlorate son descharge, and hydrogen from the cathode The impure gas obviously requires a higher pressure or lower temperature than the pure gas for liquefaction end some chlorine must be lost. Moreover, a limit is set to the extent of liquefaction by the hydrogen content of the

reach the explorive limit.

Liquid chlorine was first produced commercially by Knietsch for the Badische Anilin und Soda Febrik in 1888 (B P. 13070, 1888). The liquefection was eccomplished by compression with condensation at ordinary temperatures The compressor was of the liquid piston type using concentrated sulphume erid to confine and compress the chlorine gas, the the sulphuric acid was in turn moved by a hound piston of paraffin oil. The adoption of such a complicated apparatus is come measure of the difficulties met in attempting to provide com mercial liquid chlorine. Although steel is relatively unaffected by dry chloring at ordinery temperatures, at higher temperatures the rete of reaction incresses end becomes very appreciable at 130°C. This etrictly limits the com-pression ratio that can be used. Moreover, labrication is difficult, since all the usual lubri eenta are ettacked by chlorine Sulphuric acid appears to be the favourite choice as a lubricant.

At present liquid chlorine is produced by (i) compression with condensation et ordinary temperatures, (11) refrigeration alone, or (111) a combination of compression and refrigeration

Transport - The transport of liquid chlorine is companies and of authorities such as the Board of Trade for shipments by sea.

Regulations in the United States (and by voluntary concurrence in Canada) are issued by the Inter State Commerce Commission to which the Chlorine Institute (an association of pro-

ducers) acts in an advisory capacity,

In Great Britain advisory bodies appointed by the Home Office have published two relevant With olefins, aret.lenes, and bennen (ander appetrs—the Fourth Report of the Gas C. Inders creature conditions) chlorine combines directly, Research Committee (Cylinders for Laquefiable forming additive compounds The uncontrolled Gases), 1929, 11MSO, and the Welded Creation with actilene as explosive With Containers Report, 1931, IMSO, A summary tury-atune, the reaction is so violent that the of our distance with grounds and of carbon and laidness with grounds and Cee of laidness challenges of the warms anatomal regulations and of laidness challenges of laidness challenges of laidness challenges of laidness of laidness of the laid of SCL 1995, 55, 242–252, 269–272). types of container:

(a) Cylinders varying in capacity to 150 lb.

U.S.A. Eagland. Germany. (b) Drums.

1,900 lb. 2,000 lb. 1.000 kilos. capacity

(c) Tank

wagons, 14 tons 16 and 30 14 tonnes capacity short tons

In charging liquefied gas containers special precautions must be taken to ensure that ample " free space " is left so that the container cannot become completely filled with liquid with consequent risk of extremely high hydraulic pressure. The British filling ratios of 1.25 and 1.19 lb. of chlorine per lb. of water capacity for temperate and tropical conditions respectively are based on 6% free space at 45°C. and 65°C. These ratios are for solid drawn cylinders; for welded containers the British filling ratios

are 1.24 and 1.16 based on 6% free space at 50°C. and 75°C. respectively.

Containers should be stored in a cool, dry place, preferably fireproof. Care should be taken in the handling of containers; the strength of the containers should not be abused by exposing them to falls. If a container is allowed to remain coupled to an absorption plant after all the liquid chlorine has evaporated, the residual chlorine in the pipe-line and container will dissolve, causing the liquid in the absorption vessel to be drawn back into the container. This might have serious results, owing to the decomposition of some of the liquids under certain conditions causing a burst.

Applications.-The largest consumption of liquid chlorine takes place in the paper industry, where it is used both for the preparation and

bleaching of pulp.

In the textile industry it is used in the bleaching of cotton and viscose and in the production of unshrinkable woollen goods. When used for bleaching it may be absorbed in caustic alkali to make the corresponding hypochlorite, or it may be absorbed in sodium carbonate solution to make hypochlorous acid:

$$Na_2CO_3+Cl_2+H_2O$$

= $HOCl+NaHCO_3+NaCl$

This acts approximately twice as energetically as the corresponding amount of neutral hypochlerite.

It is used in the manufacture of chlorinated organic products, such as carbon tetrachloride, trichlorethylene, chloroform, monochloracetic acid, and derived products like ethylene glycol and ethylene oxide; for the chlorination of methane to methyl chloride and methylene chloride, and for the chlorination of benzene, toluene, naphthalene and diphenyl.

Chlorine is used for the preparation of the chlerides of aluminium, boron, titanium, sulphur, antimony, iron and tin, and for degassing molten aluminium prior to casting. Formerly considerable quantities of tin were

Liquid chlerine is supplied in three different process for detinning of scrap. The process is still in use, but is of less importance for tin recovery as the percentage of tin on tin scrap has steadily fallen from 2% to less than 1%. The residual iron is recovered by smelting.

Phospene is made by the combination of carbon monoxide and chlorine, and sulphuryl chloride by combination of sulphur dioxide and

chlorine.

Many of the above uses of chlorine have been developed because of the necessity of finding new outlets for the enormous quantities of chlorine produced in recent years by the electrolytic processes for preparing eaustic potash and caustic soda.

Certain organic chlorine compounds having lachrymatory, sternutatory, and other toxic. properties were used in the Great War (J.S.C.I. 1918, 37, 127 R, and Tone, Chem. Met. Eng. 1918, 19, 357) (v. CHEMICAL WARFARE).

TREATMENT OF WATER.—The complete sterili-

sation of potable water and complete or partial sterilisation of industrial waters such as condenser water for power stations, water for swimming baths, trade effluents from works (discharging fermentable matter) and sewage, has become a most important matter for the general well-being of the community.

The importance of sterile water for domestic purposes has long been realised, and the incidence of water-borne disease may be said to be completely within control since the adoption in one form or other of the application of

ehlorine in small regulated quantity

Chlorine treatment is also applied to sterilise water for swimming baths, to condenser water in order to prevent the growth of low forms of life in gelatinous form which prevents efficient transfer of heat, and to sewage and fermentable trade wastes in order to prevent undesirable anaerobic fermentation and to assist in the control of acrobic fermentation.

There are three general methods in use, all of which involve the use of instruments to provide measured quantities of chlorine. These methods are (1) chlorine treatment, (2) chloramine treat-

ment, (3) hypochlorous acid treatment.

In water or sewage, etc., to which chlorine, chloramine, or hypechlorous acid has been added, the chlorine will occur in various forms, such as free chlorine atoms and molecules in solution, as OCI' ions and as HOCI molecules. as well as in the form of various compounds with organic matter, etc., such as chloramines, and possibly also loose addition compounds. Most of these are known to have powerful germicidal properties, and the following suggestions have been put forward in explanation:

(1) That sterilisation is due to direct chlorina-

tion of the eell protein.

(2) That sterilisation is due to exidation of the cell material, the oxygen being formed by the action of the chlorine and/or chlorine compounds on water.

There is a certain amount of evidence in support of cach of these hypotheses, and it seems quite possible that at times both may be important factors.

That chlorine does act as a chlorinating agent recovered as tin tetrachleride in the Goldschmidt is known by the fact that the objectionable taste of certain waters containing organic matter 1

or by the chloramine process.

Water when treated with ammonia prior to its treatment with chlorine is found to have the advantage that objectionable tastes do not develop, whilst the sterilisation is just as good as with chlorine only. It is claimed for this process that the water is delivered sterile to the consumer owing to the greater stability of chlorsmine in solution.

The chlorine and chloramine treatments are used largely in Britain and U.S.A., whereas the hypochlorous acid treatment is of growing importance in Germany, where it has been anceesefully used for many years. It is claimed for the hypochlorous acid process that it has been particularly successful in many cases where neither chlorine nor chloramine treatments were entirely effective and that in other cases

it has given equally good results

The hypochlorous acid for this treatment is produced in atones are towers, where a measured quantity of chlorine is passed counter or cocurrent to (1) a measured quantity of sodium carbonate solution (G P 306193, 1916, Dentsche Solvay Werko A O), (2) water passing over marble (C P 632713, 1932, B P 412105, 1933, Ornstein); (3) water passing over moulded magnesia (G.P a 72332, 1936, Deutsche Solvay. Werke A G) See also "The Examination of Waters and Water Supplies," by J C Thresh, J. F. Beale, and E V Snckling, published by J. and A. Churchill, London.

Production of Chlorine.

Introduction —All manufacturing processes for chlorine involve the balancing of demands for chlorine and at least one other product.

With processes starting from hydrochloric acid, the other primary product is saltcake, (Na SO4), which was an intermediate in the production of sodium sulphide, soda ash, and caustic soda by the Leblane process

For many years the production of chlorine from by-product hydrochloric acid enabled the Leblane process to compete with the ammoniasoda process, but the introduction of direct electrolytic preparation of caustic sods and chlorine gave the asme end products and eventually displaced the Leblane process by reason of lower costs.

In recent times special conditions have been examined and two interesting proposals have been made

(1) The first deals with the aituation where there is a large local demand for sodium sulphate or sodium aulphide with no outlet for hydrochloric acid In this case electrolysis of hydrochloric acid to give hydrogen and chlorine has been proposed.

Westvaco Chlorine Products Inc. and F. S. Low have described (B P. 348792) a diaphragm cell for the electrolysis of hydrochloric acid

The hydrochloric acid solution enters the may be surmounted by over-chlorination cathode compartment, passes through the followed by dechlorination with sulphites, etc., diaphragm, and leaves the cell by way of the anode compartment. The weak acid is dechlormated by air blowlog and is then resaturated, heated, and recirculated.

The inlet and exit acid strengths are 20% and 10% respectively. Preferred temperatures are 45°-100°C., and voltages as low as 21 are claimed It is stated that the chlorine is of 99 9% purity and that carbon dioxide and oxygen

are absent

(2) The second proposal arises from the fact that caustie soda produced by electrolysis must displace caustic soda otherwise produced by causticising soda ash. The demand for chlorine has resulted in America in a large over produc-tion of canstic anda and efforts have been directed to working out economical alternative processes for the production of chlorine, Coupled with the above situation the market price of sodium nitrate has fallen so low that it is very much less profitable than formerly to manu-facture andmm nitrate from soda ash and synthetic nitric acid.

It has therefore been auggested that sodium sutrate and chlorine might be more profitably manufactured by the action of nitric sold on sodium chloride F. W. de Jahn (Chem. Met. Eng 1935, 42, 537)

In addition to the primary reactions,

NaCI+HNO3=NANO3+HCI, 3HCI+HNO3=CI2+NOCI+2H2O,

the NOCI reacts with water and air regenerating HNO, and HCI, the regenerated HCI reacts again with the HNO, so that the net result is the oxidation of the HCl by the oxigen in the

air with recovery of the HNO.

It is interesting to note that the similar Dunlop process was employed at St. Rollox Works, Glasgow, about 1850. Here the nitrosyl chlorida was decomposed by sulphuric acid, the consumption of which was, however, so large that the process was abandoned.

It as evident that the discovery of suitable resistant materials of construction will play an important part in the working of this process.

HISTORICAL.

1 The Older Processes; Oxidation .- Chlorine was first produced on a reasonably large scale for Javelle water and for bleaching powder by exidation of hydrochloric acid made by treating salt with aulphuric acid, by naturally occurring manganese dioxide—pyrolusite:

2NaCi+2H,SO,+MnO → Na,SO,+MnSO,+Cl,+2H,O

The development of the Le Blane process for alkalı gave rise to large volumes of hydrogen chloride, which were at first discharged rig a chimney stack to atmosphere. The patent of Gossage (1836) showed how the nursance so Copper as preferred as a tathode, but nuclei or created could be ahated by washing out the and monel metal may be used. The cathodo is gasma coke packed tower in a counter current sourcement of a daphragm of and resistant stream of water. The hydrochloric acid to abbestos cloth The anode is graphite. ganesc dioxide:

4HCl+MnO,→ MnCl,+Cl,+2H2O

The hy-product hydrochloric acid, of little value otherwise, gave a much cheaper chlorine than the previous process. Nevertheless, the imported manganese ore was expensive and the process was wasteful of this raw material. Weldon (1866) developed a successful recovery process which conserved manganese with great conomy. The liquor from the ehlorine stills, containing manganesc chloride, excess acid, iron chloride and other impurities, was carefully neutralised with ground limestone, and precipitated hydroxide of iron was allowed to settle out. The clear liquor was treated with about a 20% to 30% excess of milk of lime to precipitate the manganese as hydroxide. Steam heating was applied and the manganous hydroxide was oxidised hy air blowing, the final product being a calcium manganite,

CaO.2MnO.

The precipitate known as "Weldon mud" was recovered and returned to the chlorine stills.

The Weldon process, though more economical of manganese, still only showed about a 30% conversion of hydrogen chloride to chlorine, the remainder being lost as calcium chloride.

Deacon (1868) evolved a successful method of air oxidation of hydrochloric acid using copper chloride as a catalyst. Although this process gave a much better yield on the hydrochloric acid and prior condensation of hydrochloric acid was avoided, there were many difficulties, chief of which was the poisoning of the catalyst by arsenic and sulphur compounds. The product of the reaction after removal of excess hydrochloric acid and steam hy condensation was chlorine diluted with nitrogen from the air.

The process of Hasenclever (1883) for obtaining reasonably pure hydrogen chloride hy dehydrating the aqueous acid with oil of vitriol which could subsequently he reconcentrated was of benefit to the Deacon process, and rapid headway was made.

The Deacon process depends essentially on the equilibrium.

4HCl+O₂⇒2Cl₂+2H₂O₂

although it is presumed that the catalyst func-tions by decomposition of cupric chloride to chlorine and cuprous chloride, cuprous chloride is oxidised to copper oxychloride, and this in turn is converted by HCl into cupric chloride, the cycle being repeated.

2. The Modern Processes: Electrolysis.—The patent of Watt (B.P. 13755, 1851) covered a very wide range of electrolytic processes, including the electrolysis of brine to give chlorine and caustic soda. Cooke (B.P. 13620, 1851) described a cell for carrying out the method, and Stanley (B.P. S11, 1853) also described a process.

The high cost of electric energy at that time, however, ruled out these as industrial enter-

The difficulties experienced in the electrolytic

production of chlorine by reaction with man-| facturing scale have been first the cost of the electric current, which was absolutely prohibitive until the dynamo-electrical machine had heen so far developed as to he a commercial reality, viz. in 1872 by Gramme. When it hecame possible to conduct electrolytic experiments on the large scale over long periods of time, it was soon found that there were many and scrious difficulties, as the corrosion of the anodes, the evolution of gas from the cathode, the rapid disintegration of the diaphragms, which were either excessively porous or not sufficiently porous. There were also excessive resistances, voltages, and the generation of heat, and there were secondary reactions, so that neither the chlorine nor the alkali metal product was ohtained in the expected purity or quantity.

The various electrical processes suggested or tried for the decomposition of alkali chlorides are all attempts to overcome economically these difficulties. Workers in this field, such as Fitzgerald and Motley (1872), Wastchuk and Gloushoff (1879), and others continued to patent

processes.

An industrial electrolytic alkali-chlorine process may he said to date from the work of Matthes and Wcher (D.R.P. 34888, 1885), who discovered a satisfactory diaphragm hy means of which separation of anolyte and catholyte could be achieved. The diaphragm was made from Portland cement and salt, the latter after removal in solution giving the desired porosity. This achievement was developed by Chemische Fahrik Griesheim, later Griesheim-Elcktron, and the first large-seale plant taking 400 H.P. was operated in 1890.

About 1890 numcrous different types of cell were patented, of which that of Le Sueur (B.P. 5983, 1891), which was operated in the Eastern States of America, should be noted as the first horizontal diaphragm cell. Carmichael's cell (U.S.P. 518710 and 637851)

was also developed in the same area.

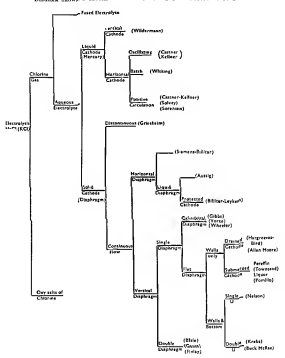
These cells, like that of Griesheim-Elektron, were discontinuously operated diaphragm cells with submerged cathode. The cathode chamber was filled at the beginning of each cycle with water or dilute caustic soda which gradually increased in strength. Diffusion of sodium from the anode chamber which necessarily took place and contamination with sodium hypochlorite and chlorate necessitated after a time cessation of electrolysis, removal of the catholyte and recommencement of the process after addition of fresh liquor.

The Hargreaves-Bird cell (B.P. 18871, 1892, and others) introduced a new principle in the non-suhmerged cathode. The cathode in this case, made of perforated steel sheet or iron wire mesh, was in close contact with the diaphragm, and the cathode liquor trickled down the cathode

and the diaphragm wall to a sump.

To reduce the tendency for hydroxyl ion to diffuse back to the analyte, carbon dioxide was admitted to the cathode chamber and sodium carbonate was produced. Steam was also admitted with the carbon dioxide to assist in heating the cell. This process was operated until 1928 by Electro-Bleach and By-Products, decomposition of alkali chlorides on a manu- Ltd., at Middlewich, Cheshire, and small units 42 CHLORINE.

DIAGRAM SHOWING CLASSIFICATION OF CHLORINE CELLS ACCORDING TO TAPE



were creeted at cellulose factories in Austria and U.S.A. This typo of cell, omitting the use of carbon dioxide, is the forerunner of three cells, viz.: Allen-Moore, Gibbs and Nelson, which, with numerous slight variants, represent the most frequently employed diaphragm cells at the present time.

The development of a horizontal diaphragm in the well-known Siemens-Billiter cell took place in 1907 about the same time as the more modern vertical diaphragm cells were evolved.

An attempt to dispense with diaphragms completely gave rise to the bell-jar or specific gravity cell, which was investigated by Richardson and Holland at St. Helens in the 'ninetics (B.P. 2297, 1890). The only industrially successful cell of this type was that of Verein fur Chemische und Metallurgische Produktion (D.R.P. 141187, 1900) installed at Aussig (Czechoslovakia).

The mercury cell was devised almost simultaneously by Castner (B.P. 16046, 1892) and Kellner (D.R.P. 70007, 1892), and others. This process, which was working industrially in 1895, has been developed from that date chiefly by the Castner-Kellner Alkali Co., Ltd., in England, the Castner Electrolytic Co. (later Mathieson Alkali Works) in U.S.A., and hy Sol-

vay & Co. in Europe.

In more recent times attention has been directed to the electrolysis of fused sodium eldoride to give sodium and chlorine, see Danneel and Stockem (Z. Elektrochem. 1905, 11, 209); Danneel, ibid. 1924 30; Gesellschaft fur Chemische Industrie in Basle (The Ciba Cell); (G.P. 236804, 1909; B.P. 18300, 1910; 17047, 1911); Roessler and Hasslacher Co. (The Downs Cell), (B.P. 238956, 1924).

The Ciha cell is worked at Monthey, and the Dewns cell is worked by the Roessler and Hasslacher Chemical Co. in America, and by Deutsche Gold- und Silber-Scheideanstalt, in

Germany.

Further reference should be made to "Monographien über angewandte Elektrochemie," W. Knapp, Halle 1933 (vols. 23, 41, 33); Chem. Trade Journal, 1924, Elliott, "Electrolytic Caustic Soda"; J.S.C.I. 1924, 43, 1224, 1250, 1291, Hale, "Modern Electrolytic Cells"; Chemical Markets, 30, 147, "U.S.A. Chlorine Supplies"; Mantell, Performance and Characteristics of Electrolytic Cells, (Chem. Met. Eng. 1931, 38, 88).

The effective development of aqueous electrolytic chlorine processes thus occurred between 1890 and 1900, and from the beginning of the century this was an established industry, rapidly expanding at the expense of the Le Blane process which died out by 1920. The industry is still growing owing to the increasing demands for chlorine products and the discovery

of new uses.

The number of types of cell in operation is quite large, as will be seen in the diagram on p. 42, which, although not complete, includes the important cells in the different classifications to which they belong, as well as some that have ceased to be used. The cells will be described under the broad classifications of diaphragm and mercury cells.

Production of Chlorine by Oxidation of Hydrochloric Acid.

(a) MANGANESE DIOXIDE.—Technical. There is divergence of opinion as to the course of the reaction between manganese dioxide and hydrochloric acid.

The first reaction is either

$$MnO_2+4HCl=MnCl_4+2H_2O$$
,

or more probably,

$$2MnO_2+8HCl=Mn_2Cl_6+4H_2O+Cl_2$$

(cf. W. W. Fischer, Chem. News, 1878, 37, 250; U. S. Pickering, *ibid.* 1879, 39, 225; Berthelot, Compt. rend. 1880, 91, 251).

The chlorides MnCl₄ and Mn₂Cl₅ form dark brown solutions which quickly decompose even at ordinary temperatures, and the ultimate

result is

$$MnO_2+4HCl=MnCl_2+2H_2O+Cl_2$$
;

but this reaction is only completed at 100°C., and with a certain excess of HCl—in practice at least 10% and more if low strength acid and hard manganese ore are used—even to twice the theoretical requirement.

The manganese chloride liquor from the stills also contains free acid, the chlorides of other metals present in the ore, and a little chlorine.

Black's analysis of still liquor in a wellconducted works showed how incompletely the acid is utilised.

> MnCl₂ . . . 10·6 HCl . . . 6·6 Al₂Cl₆ . . . 0·6 Fe₂Cl₀ . . . 0·5 Cl₂ not determined. H₂O (by diff.) . 81·7 100·0

On the assumption that all the manganese in the ore was in the form of MnO₂, and that no hydrochloric acid gas was carried away by the chlorine, it follows that out of every 100 parts of HCladded 33.6 parts had been left untouched.

The original source of manganese was invariably pyrolusite, which occurs in quantity in Germany, Bohemia, Transylvania, Russia, India, Australia, Japan, U.S.A., and scantily in North Wales.

A good pyrolusite should contain most of the manganese as dioxide, since the lower oxides are less useful.

Russian pyrolusite varies between 75% and 85% MnO₂, and averages 80% MnO₂, 1% MnO, 3% H₂O.

Impurities such as BaSO₄ and SiO₂ are objectionable as they coat particles of the oro and prevent reaction with the acid. Carbonates waste acid and add carbon dioxido to the chlorine, rendering the chlorine less suitable for making bleach. Soft pyrolusite is more easily attacked by the acid than hard pyrolusite, and is therefore preferred.

Chlorine Generation.—For a small production of chlorine, the generating vessel or still may be

made of stoneware. Fig. I shows one of the These stills measured inside 7 ft. by 5 ft. by 3 ft. best forms of stoneware still intended to be and upwards. heated by a water or steam bath.

in and taken out of the still with special tongs through the large opening. The small paper serve as acid inlet and gas outlet respectively The still liquor at the end of the operation is siphoned out or run off through a discharge pipe passing through the steam packet These small stills permit of careful and economical working

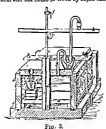




so far as materials are concerned, since the acid is not diluted by condensed steam

Fig. 2 chows a U shaped pipe of lead or earthenwere which permits of scaling a still of a bettery from the chloring main by means of a water seal-a most practical device.

For generating large quantities of chlorine the stills are made of highly silicious sandstone or other stone not acted upon by acids, impervious to them and not hable to creck by rapid changes



(4) False bottom.

(b) Hydrochloric acid inlet scaled in a bowl. (c) Lead steam pipe connected to

(d) Stoneware steam column perforeted above the bottom slab

(g) Chlorine outlet.

The small stills were worked by first adding the The manganese ore is placed in a perforated and, then inserting the perforated ore container evlinder, capacity about I cut., and this is put ready charged and rapidly making the joints as the reaction begins forthwith. When it slackens heat is gradually applied until the acid is exhausted. The still cools down overnight, the perforated container is removed and washed, and after removal of the still liquor the cycle is recommenced

The large atone stills are charged with 6 to 10 cwts of ore broken into pieces, the manhole closed and acid run in at first quickly and then gradually over several hours. The heat of reaction assists the evolution of chlorine, and when this slows down steam is injected cautiously, the temperature rising to 90°C. In the early days of the process the waste liquor evolving chlorine caused considerable nuisance and it became necessary to run it into closed vessels to which chalk or limestone dust was added, and this was subsequently the first stage of the Weldon process.

The chlorios evolved is always contaminated with HCl gas and water vapour, and these are removed by air or water cooling of a long string of earthenware or lead pipes in which the condensation of aqueous hydrochloric acid takes place and is drained away by trapped escape

Recovery of Manganese and Method of Utilisa-tion -The foundations of this process were laid in 1837 hy Gossage, who precipitated the mac-gacese with lime, ran off the supernatant liquor, and agitated the precipitate with air; oxidation was, however, very slow, and was never even half complete In 1866 Walter Weldon made his first attempts and by 1869 his new process was in commercial use at the works of Mesars. Gemble & Co, St. Helens, and by 1871-73 had been so improved as to be adopted by all British and Continental alkali works
The apparatus employed bas received few

modifications, but the conduct of the process which in the critical part is complicated and still not thoroughly explained has been more modified. In the following the plant, Fig. 4,

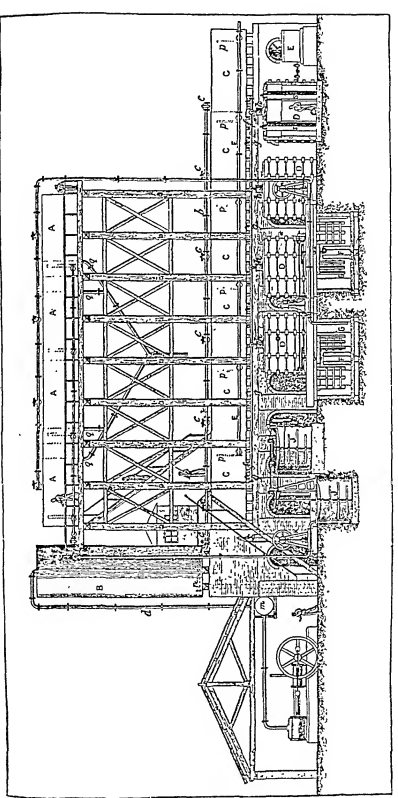
and process are described

The process is commenced in the stone still E. The only changes demanded by the Weldon process are (1) that the acid employed should be as free as possible from sulphuric acid, since CaSO, 2H,O, subsequently formed entails extra expense in filter pressing; (2) that the run off pipe for the still liquor abould lead mto a mud still D, which is shown in section in Fug 4

The neutralising wells G, G are sunk in the ground on a bed of puddled clay, and after being built are currounded by puddled clay to present possibility of leakage; they are built of large ctone clabs of acid-proof Yorkshire flag On the top of the wells two large timber Charging opening not shown
Charging opening not shown
Charging opening not shown
Such stone, found at Southon

Such stone, found at Southon

Such stone, found at Southon The best of the state of the st



TG. 4.

the wells with strong planks well tarred and by leading the gases to the suction of the nearest chimney hy an earthenware pipe. When the free acid has been neutralised yielding calcium chloride and Fe and Al chlorides are converted into the respectivo hydroxides, these last carry down as orsenites any arsenious acid originally present in the hydrochloric acid. Any aulphuric acid present in the hydrochlorie acid is converted into CaSO, 2H.O. plete neutralisation is necessary to remove the impurities and to prevent corrosion of the settling tanks A. A into which the liquor is next pumped, hut the excess of CaCOs employed should be the amallest possible to reduce the cost of the aubsequent operations

A awivel pipa is lowered from above into the hourd, as shown in Fig 4, in the left-hand well, and the turbid liquor is then piped to the gunmetal pump shown in the figure in front of the third tank D from the left. The figure shows the delivery pipe rising to the top of the plant, then running bonzontally and discharging As soon as the well G is emptied and the pump stopped, a small valve is opened at the base of the pump delivery pipe, the pipes drained back into the well, and the swivel pipe is with drawn, to prevent corrosion The well is then ready to receive its next charge

The chloride of manganese ecttlers are made of in thick boiler plates The processes already described must be repeated until at least one settling tank A is quite full. After making the last addition the liquid should settle quite clear within a few bours, otherwise the charge in the next operation is apt to froth over, and also the rerovered mangeness will consume an undue amount of acid.

After the volume of the liquor in the settler A bas been estimated and a sample teken for the flow of the clear liquor is controlled by a cock, tower B.

The oxidising towers B are made uf in. thick boiler plates, and they may be 8 ft. diameter and 28 ft. high, the bottom is flat, and the top is quite open. The pipe bringing the manganese chloride liquor enters the tower about 6 ft. from the top of the tower and the top of the swivel pipe may he set to give exactly the desired amount of liquor. When the liquor cesses to run the cocks are closed, and the reaidual liquor in the settler is reserved for the finishing of the charge.

component for the remaining processes, and analysed for MnCl, by diluting a measured carbonic acid gas, which in escaping earries sample with boiling water, adding zinc oxide away the amail quantity of dissolved chlorine until the mixture is white as milk, and, lastly, gas; these two objectionable gases are prevented adding standard potassium permanganate with from escaping freely into the air by covering occasional boilings, until the supernatant clear the wells with strong planks well tarred and by liquor ahows a pink flush; the reaction yields a manganito exactly like the Weldon process under consideration :

ZnO+MnO+O=ZnO MnO.:

and the necessary exygen is obtained from the permanganate similarly:

3ZnO+2KMnO =K.O.ZnO+2(ZnO·MnO.)+30

The temperature of the manganese liquor in the tower will be above otmospheric temperature, and in subsequent operations will be still higher, but experience has shown that if it is below 55°C, it abould be heated to this point by blowing in steam. The steam pipe is not shown in the figure, and because of the shaking of the tower it must pass over the top of the tower, and terminate at the bottom of tha tower in several branches containing perforations

The manganese chloride liquor during the then running boutontally and discharging above meaning may no agreement through control cocks into one or other of the sequent operation must be well agitated. This agitation is effected by a blast of air. used for agitating purposes alone the blast of air required is not great, but the blast can be regulated as desired, and can be increased very considerably.

Milk of hims is prepared in the upper of the two tanks J, J shown in Fig. 4. Each of these tanks is provided with an agitator. The upper tank only contains a lime cage, the bottom of which is indicated by a horizontal line above the third agitator blade. This tank is filled with water, while steam is also blown in, until the bot water stands well above the level of the hottom of the cage. Quicklime is selected so as to contain less than 1% of magnesia; it should be well hurnt, but not over burnt. The magnesia se very objectionable, because it accumulates during the repetition of the following processes, analysis, the clear liquor may be drawn off from and in the generation of the chlorine it neutralises analysis, the creat sequer may be trained on avoir same trong generation where continues to extend the borning a saved pipe just under lead to no parpose. The celected quickhine in the surface of the luque, which can be seen to be, the form of large lumps is put into the cape, quite clear. The bottom end of the sarred is some slakes, generating thereby much appear in a statched to the sade of the tank about a pipe is attached to the sade of the tank about a dark and thus addition is continued that the most bottom of the tank as as to have registers 50 on Twaddfull hydrometr, the plenty of room for the mud to collect and to be test jar being gently rotated the while; the atored pending its removal. Ontsido tho tank milk then contains 20 lb. CaO per cu. ft., and is as ofreng as can be conveniently prepared and cocks control its entrance into the unidising. The milk is then run from a valve at the bottom of the tank on to a wire sieve (not shown in tha figure), and thence auto the hottom taul J, which is used as a combined store and measuring tank.

The next operation is the addition of milk of lime to the hot agitated manganese calcium chloride liquer in B It is possible to calculate the amount of milk of lime required to (1) theoretically precipitate all the MnCl₃, and (2) increase the total amount to 16 times the theoretical. Long experience with the process and due appreciation of opposing considerations The manganese chloride liquor should be have shown that this figure gives the best

results for ensuring the rapidity and completeness of the oxidation, the economy of lime and of hydrochloric acid, the settling quality of the recovered manganese, and the avoidance of

'stiff batches.''

The milk of lime is pumped into the tower B by the pipes shown. During this operation the utmest vigilanco is required in testing, samples being drawn, filtered, and tested as rapidly as possible. The first test is simply that for alkalinity with red litmus paper. A second test is made by adding to the filtered sample a streng selution of bleaching powder, which produces a brown precipitate or colour if any manganese is still present, in which case either more milk of lime must be added or a little more time given for that already added to react. When just sufficient lime has been added to remeve all the manganese from the solution the addition is stopped.

The amount of lime so far added is always found to exceed that expressed by the simple

equation

$$MnCl_2+Ca(OH)_2=Mn(OH)_2+CaCl_2$$

because of the solubility of Ca(OH)2 in CaCl2 selution and because milk of lime always contains particles that react very slowly. The excess lime may vary from 0.10 to 0.45 (the theoretical lime given above being 1.00).

The amount of calcium chloride present profoundly alters the solubility (and hence the reactiveness) of the ealeium hydrate, as is shown by the following figures for 60°C.:

Percentage $CaCl_2$ in solution . 20 25 15 100 e.e. dissolve g. of CaO . . . 09 . 11 . 13 . 17 . 22 . 29 . 38

On cooling very long crystalline needles of the oxychloride ClCa·O·Ca·OH+7H2O separate and so much so from strong, hot solutions that the mixture becomes apparently solid and can be inverted, and this has been assumed to he one of the reasons for the production of the "stiff batches" mentioned.

Starting from native manganese the liquor will centain about 18% CaCl₂ only, and because of the lew solvent action of this concentration of calcium chloride on the oxeess lime already added, it is not permissible to add any further excess until some of the free lime has been removed by converting it into calcium

manganite.

For this purpose the hlast of air is increased to its maximum and continued until the end of the operation. The fixation of the oxygen causes the temperature to rise a few degrees, and the reaction $CaO + MnO + O = CaO \cdot MnO_2$ reduces the amount of free lime and changes the colour of the precipitate from light yellow to brown, and then to black. After about a quarter of an hour one-fourth of the remaining oxcess lime desired is added, and these additions repeated after three more periods of blowing.

During the progress of the blowing, filtrates

simultaneous oxidations of the mixture (CaO+ MnO) and of the excess MnO to form (CaO·MnO₂) and (MnO·MnO₂) respectively are now complete, and continued blowing has no further effect.

But by the addition of a second supply of manganese chloride liquor to the blowing towor a new reaction is now brought into play, namely

2(CaO·MnO₂)+MnCl $=(CaO\cdot 2MnO_2)+CaCl_2+MnO_1$

and hy continuing the air blast the MnO is eonverted into (MnO·MnO₂), as in the first part of the operation. The "blowing out" of the manganese becomes slower with each addition, and when it hecomes too slow no more is added, and the whole operation is ended. The effect of this last operation is to increase the amount of chlorine that can be manufactured from the mud, without increasing the amount of acid required for its manufacture.

The mud, consisting of calcium dimanganite, CaO·2MnO₂, and manganese manganite, MnO·MnO₂, suspended in a calcium chloride solution is now run from the base of the oxidising tower hy the cock and main c, c, Fig. 4, into the mud settlers C, C, similar in construction to the settlers A above. After a few hours the settling is complete and the clear supernatant calcium chloride liquor may then he run away through the main E, E, hy opening the cock near the bottom of the tank and gradually lowering the swivel pipe p, until the black mud below is

reached. The manganese mud is now ready to be used in the manufacture of chlorine. It differs from native manganese in that it is entirely soluble in hydrochloric acid and that it dissolves very much more readily. The generating tank therefore does not require a false hottom, and it is made very large. Fig. 4 shows four such stills D, D, that may be 12 ft. diameter and 12 ft. deep. They are oetagonal, but otherwise resemble in construction the nativo manganese still. The right-hand end still is shown in section, the two columns S, S are the perforated stone blocks for hlowing in steam. The outlet liquor pipes are not shown, nor are the gas outlet

pipes, nor the acid inlet pipes.

Into the still is charged any liquor available from the still E, hecause though it is not strong enough to act further on nativo manganese, it will aet on recovered manganese mud, and this is one of the advantages of the Weldon process. Hydrochlorie acid is then also run into the still till the acid liquors are 2 ft. deep. The settled mud in the settler tank C may then be stirred by a hand rake, to make it a little more fluid, and run through a cock g situated in the hottom of the settler, into the main f, f, and thence through another cock into the luted trap h, when it runs into the still. The ehlorine is given off very readily, and the rate of evolution of the ehlerine is governed by controlled addition of the mud. As the acid hecomes exhausted steam is blown in to raise the temperature, and the further additions of mud are made with eaution, until samples from samples will show a diminution and taken from the still by an carthenware cock then the disappearance of the free lime. The | (not shown), cease to he clear yellow and hecome a clear coffee colour that is not removed by interfer steaming. As supile tested with standard state of a state of a continuous cont

repeated.
When sufficient mud has accumulated in settlers A, A, a long-handled plug (indicated by a dotted line in each of that attack) is lifted, and tha mud is run through pipes q to filter presses (not shown in the figure), where as much manganese chloride liquor as is possible is recovered, and the cakes of stiff, nearly dry

mud are then thrown away.

When sufficient manginess chlorids luquor has been mads from native manganess in the still E to stock tha settlers A and C, and to give working charges in G, B, and D, then only so much instive manganess requires to be dissolved in E as will counterbalance tha unavoidabla losses in the whole of the Weldon plant,

Chira Asia Ordania - The Gasous Hydrochora Acid and Almonphere Oxygen Method— After many unsuccessful attempts made by Orland (1845), Juliuno (1846), Binks (1860 and 1852), Dufrène (1865), and others, the labours of H. Deacon and F. Hurter, from 1863 onwards, produced an entirely anocessful chlorine process which is based on the direct ovalation of the H in HCl by atmospherio oxygen, and which unities much more of the HCl than any process comprovement attented later (Henderson, 1871, Weldon, 1871, Wigg, 1873, set proceed on the same lines but have not been practically successful.

The Deacon process starts from the wellknown fact that a mutture of hydrochlora ead gas and oxygen at a temperature just below at red heat, especially in contact with poroa substances, partially changes into chlorine and water: 2HCH-O=H,O-C1.7. This decomposition is very much assisted by the employment of substances which serve as oxygencariers, and of these the saits of copper (previously employed by Yogel, 1855; Gaity and Laureas, 1860; Mallet, 1860) have been found more useful. Probably in all cases cuprachloride is formed, which at once splits up mito CuyCl, comtance with oxygen to form oxychloride, CuO CuCl, and this acts again upon HCl, yielding water and reforming cupics chloride, so that the reactions can begin over again:

(1) 2CuCl, Cu,Cl,+Cl,

(2) Cu Cl+O=Cl-Cu-O-Cu-Cl

(3) Cu.OCl.+2HCl=H.O+2CuCl.

Theoretically, all the HCl would thus be conrerted into chlorine, but in practice, under the best conditions, about two thirds of this decomposition is effected. The undecomposed HCl is, however, not lost, but is recovered and can be utilised for other purposes, eq., for the Weldon process.

has been studied by Lunga and Marmier, by Lewis (J. Amer. Chem. Soc. 1906, 28, 1380), and by von Falckenstein (Z. physikal. Chem. 1907, 59, 313). Lewis and Randail ("Thermodynamics and the Free Energy of Chemical Substanes," 1923, McGraw-Hill Book Co., Inc.) have emissally examined these results and those the control of the control of the control of 1906, 51, 369 for the hydrolysis and column of magnesium chloride and have shown that all these results are in excellent agreement with the equilibrium constants calculated from the free energies of steam and gaseous hydrogen chloride, which are, of course, obtained by entirely independent methods.

For the reaction

1C1,(g)+1H2O(g.)=HCI(g.)+1O.

they give the equation $\Delta F^{\circ} = 6835 - 0.02T \ln T - 0.00085T^{2} + 0.00000185T^{3} - 7.27T.$

where △F° is the change in free energy.

From the equations

 $K = \frac{[O_2]^{\frac{1}{2}}[HCI]}{[CI_2]^{\frac{1}{2}}[H_2O]^{\frac{1}{2}}}$ $-RT \text{ in } K = \wedge F^{\circ}$

d −RT in K≕∆F

the equilibrium constants may be calculsted. Although the equilibrium equation is in terms of activities, pressures (as atmospheres) may be used as a first approximation.

The following figures are calculated from Lewis and Randall's equation :

Temp. °C. 300 350 400 450 500 550 600 ΔF° cals. 2,350 1,940 1,525 1,110 690 270-154 K. . . 0-13 0 21 0 32 0 46 0 64 0 89 1 09

Deacon's process deals with the gaseous hydrochloric acid, as it as rovlered in decomposing common salt by sulphuric acid without condensing it into liquid acid. In practice only the gases from the decomposing pots, say 68-70% of the total HCI, are used for the Deacon process, those coming from the rosister being too dutte and containing too mach sulphure acid.

The quantity of air necessary to form chlorina

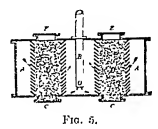
enters through the joints of the doors and dampers and is regulated by the suction maintained on the whole plant. Usually there are 4 vols. of air to 1 vol. of HCI. An excess of air is less injurious than a deficiency of it, in which cass there is a poor decomposition. The gasse leaving the pot are first cooled by a

long atung of puea and a scrubber; thus much of the water and about 1 or 4 of the HCl ar removed. Further, the gas may be dred in a sulphune acut dower before it enters the decomposer, such complete drying being very advantageons for the process. The gases now enter the heating furnace in which pipes are arranged in two sets of twelve.

The gases are here heated up to 500°; that waste heat of the furnace is sufficient for heating the decomposer, which does not possess any five of its own.

(Fig. 5), an upright, cast-iron cylinder. It contains a cylindrical ring of broken bricks D, D, supported by two cylindrical cast-iron rings of shutters placed concentrically one within the other. The gases enter at the circumference at A, pass radially across the decomposing mass D into the inner space, and are led away through the pipe OB. The decomposer temperature ought to be between 480° and 500°. The catalyst mass consists of burnt clay broken up into lumps, or ordinary bricks broken, freed from dust, and dipped into a solution of cupric chloride. The mass contains about 0.6-0.7% of metallie copper.

When the gases leave the decomposer, they consist of a mixture of HCl, free Cl₂, and steam. At best two-thirds of the HCl is converted into chlorine, but sometimes only one half. The hot exit gases are now cooled by passing through a long string of earthenware or glass pipes, and are then freed from HCl by washing with water in the ordinary acid condensers, consisting of stone towers filled with



coke, combined in some works with acid eisterns. Woulff's jars, etc. By managing this process, all the acid can be obtained in the state of concentration required for the Weldon process and other purposes.

There is theoretically no loss of HCl in the Deacon process, the whole of the acid evolved from the salt being either converted into chlorine or recovered by condensation as aqueous hydrochloric acid. In practice, of course, some losses occur, but they are not necessarily large.

During the early years of the working of the Deacon process results were disappointing and unsatisfactory. The fact that impurities accompanying the gaseous HCl were sufficient to seriously affect the process was not realised. The impurities are sulphuric and sulphurous acids, arsenious acid, water, and carbon dioxide.

The arsenious acid, derived from the As,O6 in the sulphurie acid used, forms arsenate of copper, which is even less reactive than the Cuso, formed from the sulphuric acid impurity. The presence of these impurities necessitated the frequent renewal of the decomposing material. Water is one of the products of the decomposition, and hence, when present in the inlet gases, increases the partial pressure of the products of the reaction and lowers the maximum percentage decomposition obtainable under the conditions of the process.

Carbonic acid derived from the fuel gases,

either of the salt-cake furnaces or of the decom- play an important part. You. 111.-4

The gases now pass into the decomposer posing furnace itself, was a serious obstacle to, and frequently entirely prevented, the production of strong bleaching powder. This difficulty has been overcome by greater care in the construction of the apparatus and in testing for leakage.

It will be noted from the above description that only the gaseous HCl from the salt-cake pot is dealt with by the Deacon process, and the liquid acid collected after the decomposers has to be sold as such or used in some other way,

such as by the Weldon process. Hasenelever, however (B.P. 3393, 1883), described a process for gasifying the HCl in the liquid acid by mixing with strong, hot sulphurie acid in a series of earthenware vessels through which air is blown. By this means a steady, continuous stream of gaseous HCl is evolved with an easily regulated supply of air. After passing through the series of vessels, the vitriol diluted with the water contained in the liquid hydroehloric acid has to be recon-centrated to the original sp.gr. The concentration is generally performed in long, lead, bricklined pans, heated by top heat, the fuel gases being in direct contact with the sulphuric acid. This process was used to a considerable extent, but from the large quantity of sulphurie acid to be concentrated per ton of bleaching powder, a very material addition is made to the cost of manufacture.

Other Contact Substances .- A general investigation on the mutual action of HCl and O in the presence of certain compounds of Cu, Fe, Mn and Cr was made by Lamy (Bull. Soc. chim. 1873, [ii], 20, 2). Copper salts were found to be more efficient than others, and with these the decomposition is at its maximum at about 440°.

The proposals to employ ferric chloride (Thibierge, 1855, and others) or chromic oxide (Hargreaves and Robinson, 1872) have not had any practical success.

Nickel oxide was proposed by L. Mond (B.P. 8308, 1886).

De Wilde and Reychler (B.P. 17272, 1889) proposed employment of a mixture of magnesium and manganese ellorides with magnesium sulpliate.

Production of Chlorine by Electrolysis of Alkali Chloride.

Technical. Theoretically a current of 1 amp. liberates in 1 hour g. equivalent × 3,600 96540

clement.

Simple as the electrolysis of alkali chlorides may appear from the formula, yet in aqueous solutions the first obvious difficulty arises from the activity of the alkali metal. Where no attempt is made to separate the cathode and anode products, the output of such a cell will be hypochlorite or elilorate according to the conditions of operation.

The separation of the two products naturally presents many difficulties, since factors such as temperature, electrolyte conceutration, ionic mobility, electrode materials, and current density

The effect of two of these factors is abown by the fact that the conductivity of solutions of alkalı chlorides increases with concentration

and with temperature The voltage of an electrolytic cell depends on the decomposition voltage of the salt, the length of the electrolytic path, the overvoltage at each electrode, and the current density of the cell. In the case of a diaphragm cell, resistance of the

diaphragm is also a factor. Thomson's formula may be used to calculate the approximate decomposition voltage (E) of a

 $\overline{n \times 0.2387 \times 96.540}$, where Q is the beat

of formation and n is the valency. Overvoltage is an inherent property, non calculable, of the electrods material Although

the hydrogen over voltage in eaustic solutions at graphite electrodes is very low, for constructional purposes metal cathodes are preferable and nickel with a low hydrogen over voltage is suitable, but iron with a slightly higher overvoltage, 0 08 volt (Caspari), is cheaper

Mercury is in a class apart with a very high hydrogen over voltage, 0.78 volt (Caspan), 10 volt (Taicl), and it is due to this property that sodium is discharged preferentially on electroly as in the mercury cell process and that sodium amalgam is only very slowly decomposed by water This fundamental property has made

possible the development of the mercury cell Chlorine over-voltage also varies with the anode material at carbon electrodes it is low. whilst at platinum or fused magnetite it is

The following determinations of single electrode potentials may be noted .

OH'	ion	discharge	+0	82	٧
CI'	,,		+1		
Н.	**	10	-0		
Na.	**	**	-2		

In a hrine solution the smallest decomposition voltage is 1 23 for the electrolysis of water, but the over voltages are high and at higher

voltages little of this process proceeds At 4 08 volts electrolysis of anhydrous sodium chloride commences, but since in aqueous solu tion the exothermic reaction between sodium and water to give caustic soda and bydrogen takes place, the decomposition voltage will be reduced. It will be higher than the decomposition voltage [of hydrochloric acid, 1.78 volts, due to the fact that hydrogen ion concentration will be low at an electrode where there is a prependerance of OH' ions from the caustic soda and there is a rise in the single potential for bydrogen ion discharge. The decomposition voltage will depend on the strength of the caustie soda as well as that of the brine, but is of the order of 22 volts

With a dilute sodium amalgam cathode the single potential of sodium is lowered by reason of its solubility in the amalgam, and the decom position voltage is of the order of 3 18 volts.

The decomposition voltages, therefore, of sodium chloride in saturated brine are approximately 22 volts for diaphragm cell processes and 3 2 volts for mercury cell processes.

The mobility of ions plays an important part in diaphragm cell processes. The hydroxyl ion with a potential drop of 01 volt at room temperature has a velocity of 6 48 mm. per hour compared with 2 41 mm. per hour for chloring, Migration of hydroxyl ion, therefore, must be opposed by a austable velocity of electrolyte through the disphragm if satisfactory yields are to be obtamed. It becomes clear, then, that for the same efficiency horizontal diaphragm cells will yield a stronger caustic liquor than vertical diaphragm cells

The choice of materials for electrodes has been hmsted in course of time by initial cost and by running costs The high over-voltage and high cost of platinum and magnetite anodes have eliminated these from oblorine cells and artificial graphite is now universally employed. This latter material, which is porous by reason of its method of manufacture, suffers from the fact that as porosity increases so the life of the anode decreases, It has been contended that in the pores where there is no appreciable circulation the aquenus solution becomes depleted of chloride ion and hydroxyl ion discharge occurs; this results in oxidation of the enode to carbon

dioxide and disintegration is promoted Cast magnetite (G.P. 157122 and 193367, Griesheim E lektron) was used to overcome the defects of porous carbon, and with particular advantage in the Griesheim cell, where the chlorate concentration in the analyte accentuated attack on the carbon.

More recently attempts have been made to increase the life of graphite anodes by impregnat. ing them with organic substances to reduce hag them with Grigano substances to reduce their porosity (Fringerald and Molloy, B P. 1376, 1872; Athun, U S P. 754114, 1904; da Vains, F.P. 567925, 1922, Koholyt, B.P. 221634, 1923; Pomilio, Giorn. Chim. Ind Appl 1925, 7, 63; Botel Maletra, F.P. 679810, 1928) Jaffe (Z Elektrochem. 1935, 42, 71-83) shows that impregnation causes an increase in the over-voltage of chlorine discharge,

DIAPHRAGM CELLS .- The Griesheim Gell-This was one of the earliest cells to be operated on an industrial ocale, although it has now been replaced largely by more economic units. It employs stationary electrolytes and, therefore, has not the same intensity of output, nor the sams economy in power and materials as later cells, but its advantages of robustness and simplicity well fitted it for the early days of electrolytic chlorine production. It depends essentially on Breuer's diaphragm (B.P. 19775. 1891) made by setting cement with acidified brine; after complete setting the crystallised salt can be washed out to leave a very porous block.

A number of boxes made of such diaphragm plates fixed in an angle rron frame rest in an outer steel container. These boxes make the anode compartment which is fitted with carbon anodes, whilst the steel outer vessel and steel plates hung between the hoves form the cathode. Porous pots filled with salt are suspended in the anode chambers to maintain the strength of the brine Fig. 6.

The method of operation is to fill the anods compariment with saturated hrine and the outer container with weak hone. As electrolysis

proceeds alkali accumulates in the cathode liquor until the concentration reaches 40 to 50 g. of eaustic soda per litre, when electrolysis is stopped, the caustic liquor drained off, and after refilling the cathode chamber the process is recommenced. In its long history the cell has naturally undergone many changes, higger cells were made and provision was made for maintaining a high working temperature and for collection of hydrogen.

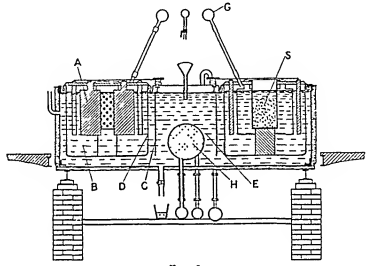
The cell voltage was of the order of 3.65, with current densities at the anodo 2 amps./dm.2, and at the cathode

0.5 amps./dm.2

Owing to the low output and absence of positive movement of solution through the diaphragm, hack diffusion of hydroxyl ion is appreciable. This leads to low efficiencies, high chlorate content, and rapid anode attack. This latter results in a high carbon dioxide content of the chlorine which in turn causes difficulty in hleaching powder production. Attempts to obviate these difficulties led to the manufacture of magnetite electrodes, but these add appreciably to power consumption owing to the much higher chlorine over voltage.

This cell was largely applied to the electrolysis

of potassium chloride in Germany.



F10. 6.

The Siemens-Billiter Cell.—The first installation of these cells was tested in 1907 at Aschersleben by the association of Kaliwerke Aschersleben, Siemens & Halske A. G., Billiter, who had carried out much experimental work previously in Vienna.

Many plants were subsequently creeted in

Germany.

The cell consists of a sheet iron tank, the cathode is a sheet of perforated iron or iron netting attached to the walls a little above the bettom of the cell.

The side walls above the cathode are covered with a layer of cement, whilst the cover of the cell consists of tiles of concreto or stoneware. The anode leads of graphite which carry graphite plates pass through the cover, whilst the iron shell is connected to the cathode leads. chlorine outlet in the cover is connected to an carthenware main, whilst an outlet pipe of iron from the cathode chamber serves to earry away the hydrogen. The caustic liquor drains from the eathede chamber continuously. Fig 7.

The cell is the only type in large scale commercial production which has a horizontal diaphragm and the success depends on having a suitable diaphragm which has sufficiently fine capillaries and at the same time a readily In actual construction the tank may be regulated perosity. The essence of Billiter's shallow, little deeper than the cathode compart-

diaphragm is the use of a fibrous material. ashestos, and a powder, harytes, which are spread uniformly on a sheet of ashestos cloth resting on the eathode. With a suitable diaphragm there is no tendency for hydrogen to penetrate, and the avoidance of this eliminates the difficulty of producing a satisfactory chlorine as well as a caustic liquor of constant and suitable composition.

The cell having been filled with brine, the circuit can be closed. The brine percolates uniformly through the diaphragm, and as the anodes practically cover the whole diaphragm a uniform current-distribution and brine-flow

A level gauge is fitted to the side of the cell to indicate the depth of brine, but a modification has been introduced in recent years. A small vessel external to the cell is connected to it just above the diaphragm and the hydrogen from the cathode chamber bubbles through the brino in this vessel. By varying the depth of the hydrogen seal in the brine the rate of brine percolation through the diaphragm is controlled at the desired rate under varying conditions of hydrogen and chlorine pressure and of brine level.

ment, and the wall of the cell may be built up] of brick or stone suitably tied together.

The cell is operated to give a constant alkalimity which is achieved by varying the level of brine in the cell. The rate of percolation and the cell voltage remain practically constant for many months, but gradually the diaphragm becomes blocked owing to the accumulation in it of calcium and magnessum deposits, and the cell is then cut out for cleaning. The depth of brme, which may be 8 cm, at the commencement of the cell life, will be gradually increased during the run, possibly to 24 cm, when cleaning is due.

The results of a run of two months with brine not specially purified are given by Billiter.

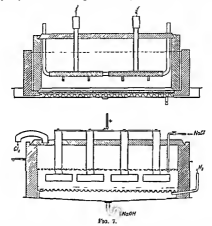
2001-B signs. Mean current . efficiency

94.7% 3 66 v. voltage .

alkalimity 130-2 g. NaOH/litre. CO, content of chlorine 1-17%

The average voltage for different current densities of a cell working at 85°-95°C. is:

Current density amps./dm. 4 6 70 Voltage 35 4.7



The alkalı content of the cathode liquor is 12- | narrow, are made of sheet iron coated anside 16% for NaOH or 18-20% for KOH with average current efficiency of 95%.

The Bell-sar (Aussia) Cell. Fig. 8 .- This cell. formerly operated by the Verem für Chemische und Metallurgische Produktion at Aussig, Czechoalovakia, is sometimes known as a gravity cell, It may be described as a disphragm cell which dispenses with a solid disphragm, since it functions by reason of stratification of the liquors The density of the caustic liquor produced at the cathodes outside the bell-jar is alightly greater than that of the feed brane, and the fluw of brine from the anode in the bell to the cathode counteracts the tendency for alkalı to diffuse back to the anode.

In a concrete vessel 8 to 10 ft long, 4 ft, wide, there rest 25 bells; the bells, which are long and

with cement, and rest on a ledge in the concrete wall. The graphite plate anodes lie horizontally inside and just above the lip of the bell



Frg. 8.

and the brane feed is distributed uniformly above the auode. The sheet iron bell acts as cathode and the caustic liquor overflows continuously. The cell is reputed to have a long life. Each

from 3.7 to 4.2 during a run, and the cell has a current efficiency of 85-90% when producing a caustic liquor containing 120-140 g. NaOH

per litro.

The Billiter-Leykam Cell. This cell was dovised as an improved hell-jar cell. cathode is placed below the bell and consists of T-section iron bars which are enclosed in tubes of ashestos cloth. The cloth is not an electrolytic diaphragm, since its purpose is solely to keep the bydrogen separate and carry it from the cell so that the chlorine may not be contaminated and the caustie liquor may not be agitated.

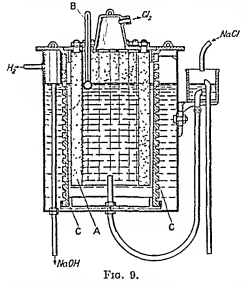
At 85°C, the voltage is 3.2, caustic liquor 150 g. NaOH per litre with current efficiency

92-94%.

The cell was intended by Billiter as a trouble-

free unit for small operators.

The Gibbs Cell.—This eell dates from about the same time as the Siemens-Billiter and has found



its principal applications in America. It was followed by the Vorce and Wheeler cells, which are derived cells different in detail.

The diaphragm here is vertical and cylindrical (Fig. 9). The eathodo is a perforated steel cylinder closed at the bottom and top with a cement diso and eover respectively. diaphragm of ashestos paper rests insido the cathode and the anodes of carbon rods inside the shell are suspended from the cover. The cell rests inside a east-iron or steel cylindrical tank,

this forming the eathode compartment.

The original patent indicated the cathode compartment flooded with caustie liquor; this method of working was ahandoned in favour of a drained cathode, which is reminiscent of the much earlier Hargreaves-Bird cell.

The Hargreares-Bird Cell .- This cell, like the

bell takes 20 amps. at a voltage, increasing | diaphragm and drained eathode which has achioved greatest success.

It was operated by Electro-Bleach & By-Products, Ltd., at Middlewich, England, until 1928, when the works was closed down.

The cell consists of a cast-iron container, 7 ft. high, 11 ft. long, 12 ft. wide, lined with cement or brick. It is divided into three compartments, by asbestos cement composition diaphragms fixed to copper mesh cathodes; the anode compartment is between the two cathode compartments. Fig. 10.

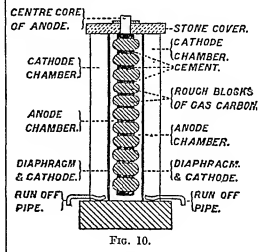
Although caustic soda was made in this cell, in practice steam and carbon dioxide were blown into the cathodo compartment and

sodium carbonate was produced.

The later cells of this type adopted newer materials and improved methods of construction

which gave a greatly increased output.

The Allen-Moore Cell.—This, like its fore-runner, the Hargreaves-Bird cell, is rectangular with vertical diaphragms. A concrete monolith makes the bottom, top, and ends of the cell, the sides being closed by shallow steel boxes



which bolt on to the concrete. The cell thus formed contains the perforated steel shoot cathode on which rests the asbestos diaphragm.

The graphite anodes pass through the concrete top, and the asbestos diaphragms inside the steel boxes holted to the concrete form the anode chamber. Fig. 11.

The standard unit takes 1,200 amps. at an averago voltago of 3.55. The eaustic liquor, 80-100 g. NaOH per litre, is produced with a current efficiency of 25%. The diaphragm life is 3 months and anode life 18 months.

The cell is principally used in paper mills in

The Townsend Cell .- This is similar to the Allen-Moore cell. A now idea is introduced in the flooding of the cathode chamber with kerosene. This compensates the hydrostatic head of brine on the opposite side of the dia-Griesheim cell, is interesting historically as phragm and reduces the rate of percolation. being one of the first to achieve commercial The production of a stronger caustic liquor is success. It is particularly interesting as the progenitor of the continuous type with vertical OH' ions into the analyte having quite the same force as if the cathode were submerged altered as required to control the rate of percelain cautic higher complete). The cautic higher tion. A 3,000 amp. cell at an average voltage is carried away from the cathode by the inf 40 produces a caustic higher higher the phylogon, and analy strough the kerosene and NaOH per titre. hydrogen, and sinks through the kerosene and runs away. Fig 12.

A standard 2,500 amp cell as 8 ft. lung, 1 ft. wide, and 3 ft deep The caustic honnr, 150-180 g per lite, is produced at a current efficiency of 95%, the cell working at 4 0 to 4 2

The diaphragm of asbestus cluth is painted with a mixture of asbestos fibre, ferrie nxide, and rolloidal ferrie bydroxide

This cell is operated by the Hooker Electro chemical Company at Niagara Falls

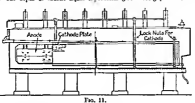
The Pomilio Cell -This cell is very similar to the Townsend and other cells in this type the cathode. The depth of caustie honnr is containing 90 tn 100 g per litre,

The first installation was made in 1921 at

Naples.

The Nelson Cell -This is a diaphraem cell concisting of a rectangular steel box in which a U shaped perforated steel sheet is attached to the tup; the ends of the channel are closed by the ends of the steel box which must, thereface, be protected inside the anode compartment by cement or conceete. The disphragm consists of aheets of asbestos paper and the graphite anodes, 14 tn the cell, are blocks 4 in by 4 m by 17 m

A cell normally works at 1,000 amps. with a It operates with a controlled submergence of voltage of 375, and produces eaustic liquor



the Edgewood Arsenal plant of the U.S. Gavernment

The Krebs Cell -This is an improved multiple Nelson cell with two or more U chapes in the diaphragm The double U diaphragm cell takes 2,000 amps Means are provided for increasing the depth of electrolyte as the rate of percolation tends in decrease. At 90% efficiency cathode hours averaging 110 g NaOH per litre is produced.

Several installations of this cell have been made in Europe

The Basle Cell (Ciba, Monthey).-Known by aeveral names, this is the cell of the Society of Chemical Industry in Basie and is operated at Monthey in Switzerland and in several installations in Italy and France It is a cell of novel content nf the chlorine, and good efficiency.

The cathode is a cylindrical iron wice cage of amall diameter, 24 in long, nn which the asbestos diaphragm is deposited externally. The bottom of the cylinder is closed whilst there is a restricted nutlet, of 1-in I.D. iron pipe, so that a froth of bydrogen and caustic liquor is carried upwards into a closed channel in which they can separate.

The anode is a carbon plate which is aurrounded by an asbestos cloth hanging from the

This cell was adopted for what was at one is drilled and screwed at one end to receive the time the largestalkali chlorine plant in the world, anode current lead which is a carbon rod 11 in. diameter. A cathode unit consists of twelve cathodes

attached to the closed channel and the snode unit consists of ten anode plates attached to the cell cover, an inverted earthenware trough, which serves to collect the chlorine.

Cathode units alternate with anode units in the cell container, individual cells carrying 6,000 ampeces.

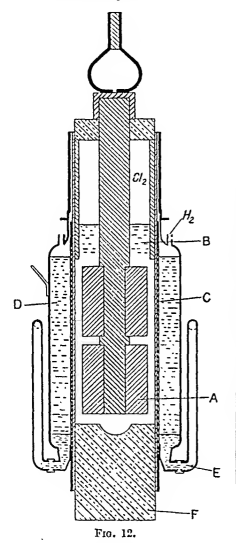
The reported performance is current density 4 5 amps /dm. at the anode and 9 0 amps /dm. at the cathode, starting voltage 3 3 v. rising to 45 v., caustic liquor 110-130 g. per litre with current efficiency of 93%.

The Gauss Cell (Badische Anilm- und Soda-Fabrik) -Little, beyond the patent specification, has been published regarding this cell, design and possesses two diaphragms which it is but it is ceported in be worked in a large acale, claimed give long anode life, line hydrogen Similar in the Basic cell in having twn diaphragms, it is, however, so arranged that the hrme feed is divided, and a portion of the main feed to the anode enmpartment is bled off to the space between the disphragms.

The pheet of this arrangement is to reduce the back diffusion of hydroxyl ion to the anode compartment, with corresponding diminution of the conversion of dissolved chlorine to bypochlorite.

The Finlay Cell .- In this double diaphragm cell rover. The plate, 24 in. by 4 in. by 2 m., cell the brane feed is introduced between the diaphragms. Successful operation depends on reducing the electrolyte gap which was achieved by a sort of filter press arrangement. The tendency for the anolyte to become denuded of chlorine ion would appear to be compensated by resaturation of the effluent brine.

MERCURY CELLS.—The Castner Cell.—This cell shares with the Griesheim, Le Sueur, and Hargreaves-Bird cells the position of being one of the first cells to prove successful on the

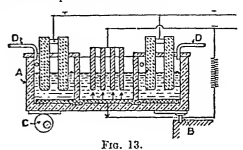


industrial scale. It is described as consisting of a box, A, 4 ft. square, 6 in. high, with two vertical partitions which divide it into three equal compartments. The whole is constructed of slate or concrete. There is a gap under the partitions of $\frac{1}{16}$ in. and a corresponding chauncl in the cell bottom which has been filled with mercury effectively seals the compartments but allows a flow of mercury. Fig. 13.

Foerster gives the cell performance as 630 amps. at 4.3 volts.

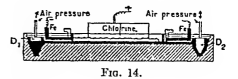
In the two outer brine compartments eblorine is ovolved, whilst in the inner compartment sodium amalgam is decomposed and caustic soda produced.

Mercury circulation is effected by oscillation, one end of the cell resting on a fixed support, B, tho other being attached to an eccentric, C, which by its rotation imparts a rise and fall, with an amplitude of half an inch.



The anodes consist of graphite rods which enter through the side of the cell, and the original proposal for the cathode was an iron grid suspended in the caustic soda. This arrangement, which makes the mercury a bipolar electrode, was never worked, since for continuous operation each side of the bipolar electrode must work with the same efficiency. Castner, in the attempt to reduce power consumption to a minimum by using this arrangement, balanced the loss in the brine cell by a regulated shunt between the iron cathode and the mercury. Finally, however, the mercury was made the cathode, and the discharge of sodium from the amalgam was achieved as in the Kellner cell by making the caustic compartment into a short-circuited primary cell with an iron cathode.

The Kellner Cell.—The construction of this cell in three compartments is similar to that used by Castner except that the cell is fixed and the mercury is caused to circulate to and fro by means of air pumps. Compressed air blown alternately into iron hells, D_1 and D_2 , immersed in mercury troughs outside each end of the cell



effects the regulated displacement and flow of mercury.

The centre compartment serves for the decomposition of brine, whilst coustic soda is produced in the two outer compartments.

The cell is made of concrete 12 ft. long by 3 ft. 6 in. wide, and the anode chamber formed by slate partitions is 5 ft. 6 in. long. The anodes of platinum wire net, 88 in number, each weighing 1 g., are mounted in concrete. The caustic soda is produced in the outer compartments using cast-iron grids as the cathodes of a short-circuited primary cell. Fig. 14.

The cell takes some 4,000 amps at 5 volts 90-85%, a 20% canstic liquor is produced ving a cathode current density of 25 amps / (Whiting, J.S.C.I. 1910, 29, 436).

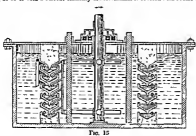
"Working at 50°C. there is a current The Wilderman Cell—This, cell has been giving a cathode current density of 25 amps / dm. Working at 50°C. there is a current efficiency of 92% and the caustie liquor contains about 30% NaOH.

The cell is operated by the Bosmsche Elcktrizitats A. G . at Jarce, in Jugoslavia.

The Whiting Cell -A small installation of this cell has been operated by the Oxford Paper Company, U.S.A. (U S.P. 951228, 951229 of 1910)

A shallow cement box is divided into five compartments by glass partitions to form the cell proper, As electrolysis proceeds, the amalgam of each compartment in turn is dis charged into a single caustic compartment where as possible to the ring channels are vertical it follows a serpentine path down channels in graphite anodes. The mercury is agitated by graphite blocks in contact with caustie liquor After removal of the sodium the mercury is pass between the anodes, the arms being elevated from the sump to a reservoir which attached to a central shaft which is actuated to recharges the brine compartments in turn.

operated in Germany. Its outstanding feature is the quasi-vertical arrangement of the mercury eathode. To provide this a series of Y section ring shaped channels are mounted one above the other and mercury is filled into each trough so that the lower leg of the Y section dips into the mercury in the ring below. These rings of iron are covered with abonite. The system of ring channels makes a cylinder which is mounted m a container. The cylinder is the brine compartment and the annulus the caustic soda compartment. Inside the cylinder and as close means of teeth at the end of arms which give rapid vibration. The whole of this Working at 40°C, with a current efficiency of mechanism is covered with abonite. Fig. 15.



sodium from the amalgam A 6,000 amp cell covers an area of 32 sq ft and works at 5 volts with cathodic current density of 20 stops /dm * at a temperature of

50°C.; 25% caustic soda liquor is produced The Kellner-Solvay Cell - The form in which the increury cell has found widest application is the Kellner Solvay cell It is operated in several slightly differing forms, chiefly in Great Britain and on the Continent

The cell consists of two concrete troughs side by sale, each with a slight slope in opposite directions, covered when working with a sheet of mercury or amalzani

In the lime compartment electrolysis takes place between either platinum or graphite anodes and the mercury rathode. Chlorine is anodes and the mereury cathode removed through a suitable exit and sodium amalgain is discharged into the caustie compart ment where it is decomposed by a counter current of water with liberation of hydrogen Fig. 16.

The mercury , stripped of its sodium, is elevated annum.

Graphite rods floating in the amalgam in the | back to the brine compartment, to begin the caustic compartment assist in discharging the cycle afresh, by means of a scoop wheel (Solvay), Archimedean screw (Castner Kellner), or vertical screw (Aussig) (Taussig, Z. Elektrochem. 1909,

15, 659). A circulation of brine is maintained between the cells and the salt saturators.

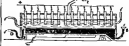


Fig. 16.

According to Foerster, this cell consists of a cement lined iron container 14 in long by 50 em. wide which, according to Billiter, takes 12,000 amps, at 5 volts using platinum anodes. The loss of platinum is given as about 11% per

centains 99.6% chlorinc, 0.2% hydrogen, and 0.1% oxygen, whereas using carbon anodes it contains 96% chlorine, 3% hydrogen and 1% carben dioxide.

The chief advantage of the mercury process is the higher concentration of the caustie liquor of greater purity. The disadvantages are tho higher power consumption, the large amount of capital invested in mercury, and the high per-

centage of hydrogen in the chlorine.

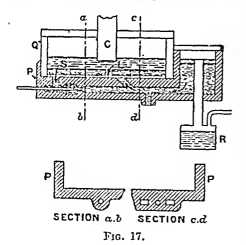
Whilst the mereury process is used almost exclusively in the production of caustic soda, it may be used for the production of caustic potash, but with greater difficulty, see Allmand and Ellingham, "Applied Electrochemistry," 1924; Kurt Arndt, "Technische Elektro-chemie," 1929; Engelhardt, "Handbuch der Technischen Elektroehemie," 1931; Foerster, "Elektrochemie wässeriger Lösungen," 1922; Billiter, "Die elektrolytische Alkaliehloridzerlegung mit starren Metallkathoden, i, 1912, ii, 1913"; Lucion, "Die elektrolytische Alkali-

chloridzerlegung mit slüssigen Katboden," 1906.

Cells with Fused Electrolyte.—The Vautin
cell (B.P. 13568, 1893, and 9878, 1894) electrolysed fused NaCl or KCl, with or witbout the addition of fluorides or other chlorides to lower the point of fusion, over a cathode of molten lead or tin with which the sodium or potassium formed an alloy (see paper by Vautin,

J.S.C.I. 1894, 18, 448).

The Acker cell is designed on the same principle as the Vautin cell, and makes use of the lead cathode (see B.P. 6636, 6637, 1898). Below is a sketch (Fig. 17) of the Acker



cell, taken from a paper by J. B. Kershaw in Electrician, 1902, 48, 20. The bottom of the cell P is trough-shaped and of east iron. It serves as electrical connection between the main conductor and the mass of fused lead L resting on it. The upper part of the cell Q is constructed of any refractory basic material not attacked by chlorine gas or fused salt. The anode C is of earbon and dips into the fused salt. The channels and subsidiary vessel R are arrangements designed to remove and decom-

The chlorine gas, using platinum anodes, pose the lead alloy as it becomes sufficiently saturated with sodium. The decomposition of the alloy is effected by blowing steam into it, and the heat generated by the conversion of the sodium into hydroxide is returned to the cell and assists in maintaining the necessary temperature. The voltage actually taken by the Acker cell is 6.75, and the current efficiency 54% ("Report of Commission on Science and Arts," J. Franklin Inst. 1903, 156, [3] 221-4). The Acker cell was worked at Niagara Falls on an industrial scale from about 1900 to 1907, when the works were completely burned down and have not been rebuilt. Fig. 19.

The Ciba cell (Fig. 18) is used for the production of sodium by the electrolysis of fused sodium chloride, the production of chlorine

being incidental thereto.

It is described by Gesellschaft für Chemischo Industric (B.P. 18300, 1910, and 17047, 1911) as constructed of fireproof brick so arranged that the chambers for collecting the products of electrolysis are separated from each other by walls or diaphragms which are mainly vertical and are capable of resisting attack by the electrolyte and the products of electrolysis.

The salt, which may be molten alkali chloride or mixtures thereof with other salts, is fused by an arrangement of heating resistances and additional electrodes for melting the electrolyte by alternating current, the resistances being

used to produce a layer of molten salt.

The Downs cell for the production of alkali mctals and chlorine is described by the Roessler and Hasslacher Chemical Company of New York (B.P. 238956, 1924) as one in which the metal halide salt is first heated, fused, and thereby freed from moisture in a large, separato compartment of the cell before it is subjected to electrolysis in the second compartment which is the cell proper.

The anode is prefcrably graphite, the cathode is either iron or copper, and these are sub-stantially concentric, whilst the cell container is a steel shell lined with suitable refractory

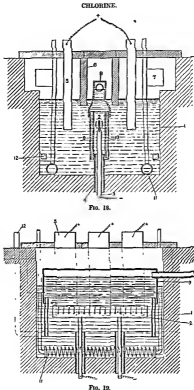
hricks.

The electrolytic compartment is formed by a combination of the sodium collector and the chlorine collector from which two concentrie diaphragms are suspended. The collector combination is submerged in the molten electrolyte and the sodium flows continuously from the collector up a riser pipe to a receiver. The upper portion of the ehlorine collector is of reduced cross section, so that above the collectors a considerable volume of molten salt is provided by the fusion compartment, and by this means access of moisture or gascous impurity into the cell proper is avoided. Fig. 20.
See also Billiter, "Schmelzfluss-Elektrolyse"

Knapp, 1932.

ELECTROLYSIS OF ZINC CHLORIDE.—The waste calcium chloride from the ammonia soda process was utilised from 1897 for some years hy Brunner, Mond & Co., of Winnington, as a source of chlorine. The process has now been discontinued. By heating the liquor with crude zine oxide and earbon dioxide the reaction

CaCl₂+ZnO+CO₂=ZnCl₂+CaCO₃



- 1. Fireproof brickwork.
 2. Cathode.
 3. Cathode conductor.
 4. Insulating sleeve.
 5. Anode.
 6. Daphragm resistant to chlorine
- 7. Chlorine ducts,
- 8. and 10. Diaphragm resistant to sodium.
- 9. Bell collector for sodium with outlet 13.
- 11. Resistor heaters.
- 12. A.C. electrodes,

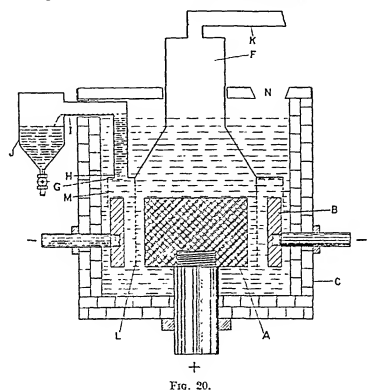
carbonate (calamine) the reaction

$$CaCl_2+ZnCO_3=ZnCl_2+CaCO_3$$
.

The filtered solution of zinc chloride is freed from its impurities (amongst which is a not inconsiderable amount of cadmium), and the resultant solution is electrolysed with carbon anodes and circular revolving eathodes that are only partially immersed, to produce chlorine gas and an exceedingly pure metallie zine. Many of the details are due to C. Hopfner (B.P. 11724,

takes place, or by treating it with native zine | xvi., "Die Darstellung des Zinks auf elektrolytischem Wege," Günther, 1904, pp. 99-180, and xxviii. "Die elektrochemische und elektrometallurgische Industrie Grossbritanniens," Kershaw, 1907, p. 108; and also in "Applied Electro-Chemistry," Allmand, 1912, p. 286.

PROPOSALS AND LESS IMPORTANT METHODS FOR THE MANUFACTURE OF CHLORINE .- Many processes have been proposed and tried for utilising the residual ammonium ehloride, or the residual calcium chloride of the ammonia 1894, and D.R.P. 85812, 1895). Fuller soda process, but with the exception of Höpfner's details are given in two volumes of "Monographien uber angewandte Elektrochemic," are now in use. Many other processes have



- A. Anode.
- B. Cathode.
- C. Cell container.
- F. Collector and done for chlorine.
- G. Collector for sodium.

H. Riser pipe for sodium.

I, J. Pipe and receiver for sodium.

K. Chlorine main.

L. M. Diaphragms. N. Hole in cover for salt charging.

also been tried for preparing chlorine from 11, 466; 1893, 12, 10, 63. There was also a various sources and in various ways, but have proposition from the Verein für Chemische been abandoned. Some of these old methods Industrie of Mainz (B.P. 3322, 1886). are enumerated below; fuller information may be obtained from Lunge's "Sulphurie Aeid and Alkali," 3rd ed., vol. 3.

1. Chlorine from Ammonium Chloride.—The most persistent efforts for the direct manu-

2. Chlorine from CaCl2.—Prolonged efforts for the direct recovery of the chlorine in the CaCl₂ produced in the ammonia stills of the ammonia-soda process were made by Solvay: B.P. 77 and 171 of 1877; 838 of 1880; 7258 facture of chlorine from NH,Cl were made and 7259 of 1884; 8724 of 1885; 13389 of by Mond, who took out B.P. 65, 66, 1049, 3238, of 1886; 10955 and 17273 of 1887; process (B.P. 731 of 1883); Bramley's process (B.P. 731 of 1885); Bramley's process (B.P. 8289 of 1887); Lyte and Steinhart (B.P. the Mond process are found in J.S.C.I. 1892,

Chloride of Lime or Bleaching Powder. hypochlorite mixed crystal with basic chloride, -Bleaching powder was first manufactured by Charles Tengant in 1799 at St. Rollox Works, Glasgow, the original patent being B.P. 2312, dated 30 April, 1799. The first year's output was 52 tons and the selling prire £140 per ton. During the nineteenth rentury the production of bleaching powder reached extraordinary tonnages, but for many years row the amount produced has been dechning. This is particularly notireable in North America, where the production at the present time is negligible. In Great Britain the tendency is also marked.

The reasons in the two cases are similar, to the extent that consumers of chlorine in the form of calcium hypochlorite find that technically and economically the production of these solutions is rapidly declining, many improvements in the from liquid chlorine at the place of use is prefermanufacture have been introduced. Attention able in the case of the electrolytic chlorine has been directed to (i) improvement of working industry in America, the Leat of summer rendeced production of bleaching powder a difficult and costly operation, and, in addition, the lack of stability of ordinary bleaching powder in a hot climate caused considerable losses due to deterioration of stocks both at the producing and

consuming works.

Composition -Until 1935 the composition of bleaching powder had not been established in spite of over a hundred years of experimental work, partly on account of the innate complexity of the problem and partly by reason of the differences in samples made in different ways.

From its method of preparotion it was

originally assumed to be a direct compound of chlorine and lime, hence the name " chloride of limo"; but in 1835 Balard postulated a compound or mixture of equivalent proportions of calcium hypochlorite and calcium chloride.

admixed with calcium hydroxide

Oding, 1861, without any proof, suggested the constitution Ca(OCI)CI, which was a ccepted by Lunge, Schappi, and Nael because they found that calcium chloride as such does not exist in bleaching powder, since the substance is not deliquescent nor is calcium chloride extracted from it by alcohol.

Although Kraut (Annalen, 1882, 214, 354) suggested the existence of a double sait bypochlorite chloride, the Odling constitution, Ca(OCI)CI, was generally accepted until recent years. The tendency in rerent times to discredit the Odling constitution arose from the work of O'Connor, (J.C.S. 1927, 2700), which showed the existence of basic hypochlorites in the calcium hypochlorite-calcium hydroxide-

water system.

A most important contribution on this subject has been made by Bunn, Clark and Chifford (Proc. Roy. Soc. 1935, 151, 141), who attacked the problem by phase rule methods, microscopic examination, and X-ray powder photographs. They showed that the first stage in the reaction is the formation of the basic hypochlorite the former is converted into another substance further from being finished until the whole of which appears to be a mixed crystal the chief the chlorine was absorbed. constituent of which is calcium hypochlorite.

Further chlorination resulted in partial conversion of the basic chloride into the tetrs. hydrate CaCl 4H,O, while the hypochlorite

mixed crystal persists.

The non-deliquement nature of bleaching powder and the difficulty of introducing more than 35-37% available chlorine into the solid are due to the presence of basic chloride, CaCl₂-Ca(OH), H₂O, which appears to be a very stable substance.

These results appear to leave the nature of the hypochlorite mixed crystal as the only substantial problem that has still to be solved.

MANUFACTURE OF BLEACHING POWDER .-Although the production of bleaching powder conditions and labour asving, (ii) avoidance of corresion of materials, (iii) production of a drier and therefore more stable product.

Lime -The quicklime used must be of high grade if a strong bleaching powder is required. Silica should be low, since calcium and magpesium silicates interfere with the settling of bleaching powder solutions. Iron compounds, being catalysts for the decomposition of bypo-chlorites, are objectionable. Manganese can, under certain conditions, be converted to permanganates and give rise to an undesired

purplish colour.

Hand elsking of hme has been entirely super seded by various entirely enclosed mechanical devices which save labour and prevent nuisance from escaping steam and dust.

Although the limestone must be well burned to prevent excess of mert calcium carbonate, a "soft hurned" lime is proferred for bleach

making. Moreover, since the chlorastice is catalysed by water, the slaked lime should contain an excess of 'tree' water.

Bleaching Powder Chambers.—The traditional lead chamber or Weldon chamber process is now almost if not quite obsolete. The chambers were 10-30 ft. wide, up to 120 ft. long, and with areas up to 3,000 sq. ft. The height was about 61 ft. to give reasonable head room. The floors were tiles or stone flags; the sides and top were 5 or 6 lb. sheet lead, fastened by external lead atraps on an external timber

framework. Floor joints were asphalted.

The chambers were preferably erected on pullars, 7-10 ft. high, so that the finished bleach could be packed through trsp doors in the chamber floor. The chambers were fitted with end doors of iron coated with protective paint. " Lutes " were provided for the introduction and

removal of gases

The 3-4 m. layer of slaked lime was spread by shovel and rake. For "gassing," the chambers were worked in series of four or more. The chlorine entered first a nearly finished chamber, Ca(OCI), 2Ca(OH), and the basic chloride and the unabsorbed gas passed in sucression CaCl, Ca(OH), H.O. On further chlorimation through other chambers which were further and

When a chamber was finished, the stream of Ordinary bleaching powder containing about chlorine was diverted into the next chamber. 35% available chlorine is a mixture of this The residual chlorine in the finished chamber was

removed either hy dusting in lime hy a Brock | and Minton's whizzer (B.P. 7199, 1886) or hy draughting into a freshly limed chamber. Great Britain it is illegal to open or vent to atmosphere a chamber containing more than 21 grains of chlorine per cu. ft. of air.

To overcome difficulties of penetration of the chlorine to the hottom of the lime layer, other systems of gassing were adopted involving the "turning" of the lime hy hand at a half-

way stage.

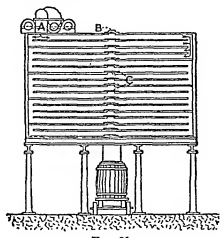
100% chlorine cannot be used without dilution, since the heat of reaction causes the temperature to rise higher than 55°, the temperature at bleaching powder decomposes which the other hand, vigorously. On \mathbf{tho} Weldon chambers are not very suitable for the absorption ef extremely dilute chlorinc. This would involve a disproportionately small output. Werked under the hest conditions the output may be reckoned as 2-2½ lb. of hleaching powder per square foot of area per day (Lunge's "Sulphuric Acid and Alkali," 3rd ed., vol. 3, p. 618; 21st Report of Chicf Inspector of Alkali Works for the year 1884).

Various plant have been devised either to absorb dilute chloring or to improve working

conditions.

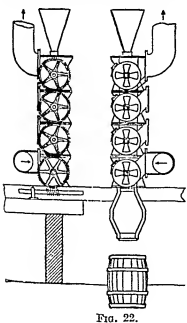
Deacen chlorine because of its low strength was absorbed in a shelf chamber plant, Fig. 21, consisting of staggered shelves which formed a zig-zag path for the gases. A shallow layer of lime was raked on each shelf, the lime heing fed through holes in the shelves which were successively covered with plates as the shelves were charged from the lowest upwards.

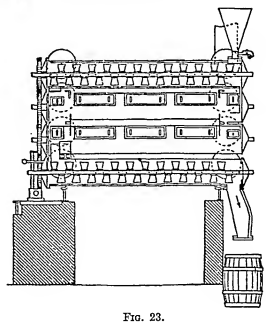
The Hasenclever plant (B.P. 17012, 1888), shown in Figs. 22 and 23, consisted of several superimposed iron cylinders fitted with central shafts carrying hlades. The shafts were driven hy geared wheels coupled together and the set of the blades caused the lime fed at the gas



Fro. 21.

outlet of the top cylinder to travel countercurrent to the ascending dilute chlorine. This plant, designed for Deacon chlorine, was used for diluted electrolytic chlorine, but like the Deacon chambers has been largely displaced by the following mechanical plant.





eight) of superimposed substantially circular the rim to the centre, hy means of ploughs on

Backman Chambers.—This apparatus invented | chambers, on the floors of which the descending by Nils Backman (G.P. 404768, 1922; 435019, limo and bleach are made to travel alternately 1925; 451114, 1926) consists of a number (usually from the centre of the floor to the rim and from cblonne. The chamber is constructed of cou-available chlorine content but was remarkable crete and independent cooling coils are embedded for its stability (see also J.S.C.I. 1921, 40, 240T; in the concrete floors. The product by this B.P. 242805, 1924, Lamble and United Alkali means can be kept at a temperature below 40 C. Co, Ltd.; B.P. 349358, 1930).

Moore Tube -The Moore rotary tube (B P. 113457, 1916, Moore and Castner-Kellner Alkah Co, Ltd) is described as a rotating exhinder not much shorter than 80 ft long, preferably with a suitable fall, and lined with tiles or other suitable material, through which hime travels counter-current to chlorine Advantages claimed are the absence from the bleaching B P. 344012, Carter and I C.I., Ltd., 1929 powder af products of corrosion of iron which cause instability, and reduced maintenance desired for use in tropical countries, which has costa

external water spray where required. Also a solvent remayed by distillation, which remayes circulatory system is adopted for the chlorine, the water at the same time. The product is gaa with permissive external cooling (see also knewn as "Sieblor" (see also knew Met. Eng. Nydegger, C P. 370841, 1920)

STABLE BLEACHING POWDER -The keeping fabriken) properties (stability) of bleaching powder have the object of study since 1874 In recent discussed by Ditz (Z. Angew. Chem. 1901, 14, years more attention has been paid to this problem with the object of producing a stable

product In JSCI 1918, 37, 311T, an article by Rettie, Lorrain Smith and Ritchia describes tests been added in order to reduce by its slaking method of comparison of the stability of bleach the free water and water of crystallization of ing powders.

cotating arms, in counter current to a stream of | bleaching powder. The product was of lower

In 1925 Conroy, Lamble, Latham and United Alkali Co, Ltd, devised a process for the production of dry and, therefore, stable bleaching powder (B.P. 246000). Further technical improvements in the production of dry bleaching powder are described in B.P. 317572. Moore, Lamble, and I.C.I., Ltd., 1928; and

Stable bleaching powder is particularly resulted in the product becoming known in the coth
Radge Tube—Tha Rudge cotary tube (BP)
126773, 1918, Rudge and United Alkah Co,
Lid jis described as a cotatable cylinder, usually
126176, 1918, respectively and context of the production of the first of the production of the first of

3, 25, 49, 105).

The fallawing table gives the composition calculated from analyses of various types of bleaching powder. Under the heading "Heat Test," are given the results of beating samples in with bleaching powder to which quicklims had test tubes to 100°C for 2 bours, a convenient

			C*OCI* C*CI*	Ca(ClO ₃) ₂	C+(OH)2	C±CO,	но	Avail Ch. after	
	able	Total							heat
Chamber bleach Haseuclever bleach Moore cotary bleach Rudge cotary bleach Backman bleach Tropical bleach Caffaro bleach	39 58 36 20 36 85 35 75 38 20 35 85 37 50	37 90	61 80 65 93 63 98 68 38 64 13		0 15 0 31 0 75 0 24 0 10 0 75 0 20	11 62 92 98 21 66 22 80 16 58 26 53 26 48	2 30 3 43 1 30 2 85 3 60 1 30 1 00	13 04 4 99 5 95 6 43 9 56 1 16 1 84	14 94 26 81 32 75 33 00

chlorine.

1141), 265,000 cals Neumann and Muller (Z. angew. Chem. 1925,

38, 194), 247,000 cals. Neumann and Hauck (Z. Elektrochem, 1926.

authorities was such that little or no water was proving mare economical. During the preparaevaporated, but no record af the free water thou of bleach liquor decomposition occurs if the content of the Fleach was given.

Bleaching Powder: Heat of Formation - scribed area. It is also made by solution of Results expressed as cals, per kg of available bleaching powder in water, the large amount of o. Nydeger (Chem Met. Eng 1923, 30, in disposal The amount of sludge may be reduced by chiarmation of the free lime, liquid chlorine being used for the purpose. As a further step, milk of lime may be directly chlorinated. Since 1918, when haud chlorine has become 32, 27), 252,000 cals.

The nature of the determination of these processes tend more and more to be adopted, processes tend more and more to be adopted. temperature exceeds 40°C. Evaporation of Bleach Liquar .- Bleach fiquor is produced figured chloring is utilised for cooling by direct in chloring works for delivery within a circum introduction of the liquid into milk of lime in

absorption of chlorine in milk of lime a peripheral speed of 500 ft. per min. for the agitation is desirable, and the depth of the ehlorinator will depend on the quantity of gas to be absorbed in unit time.

A convenient method for the preparation of bleach liquors on a small scale (batches of 70 lb. chlorine) is by the use of the Golding-Pritchard chlorinator (B.P. 233490). The chlorine valve is automatically controlled by progress of the chlorination, which causes the rise or fall of a gas bell.

In addition to the use of bleach liquor for bleaching cellulose, it is used for the removal of oxidisable sulphur bodies from the lighter

fractions of mineral oil.

Analysis of Bleach Liquors and Allied Products. The best method for the determination of available chlorine is due to Penot. A suitable measured quantity of bleach liquor or other solution of hypochilorito is titrated with sodium arsenite solution in the presence of an excess of sodium bicarbonate, the end point being determined by spotting, using starch iodide as an

The alkabinity of hypochlorite solutions is most eonveniently determined by titration of the solution with acid, using phenolphthalein and methyl orange as indicators, subsequent to destruction of the hypochlorite by means of an excess of neutral hydrogen peroxide:

$$Ca(OCI)_2+2H_2O_2=CaCI_2+2H_2O+2O_2$$

An apparatus for the rapid determination of available chlorine in weak hypochlorite liquor is described by J. M. Taylor, Ind. Chem. 1929, 5, 135.

Calcium Hypochlorite, Ca (OCI), .- Crystalline calcium hypochlorite was first prepared by Kingzett (J.C.S. 1875, 28, 404; Chem. News, 1875, 32, 22; 1882, 46, 120) by evaporation or freezing of bleaching powder solutions (see also Orton and Jones, J.C.S. 1909, 95, 751; Proc. Chem. Soc. 1909, 25, 74). It crystallises from water in minute needles with 3 mols. of water of crystallisation (Urano, Trans. Electro-chem. Sec. 1926, 49, 65; O'Connor, J.C.S. 1927, 2700). The solubility of ealcium hypo-chlorite in water at 0°C. is 21-8%. With hydrated line calcium hypochlorite forms basic hypochlorites, Ca(OCl₂)·3Ca(OH), and Ca(OCl) ·2Ca(OH), which are less calcula-Ca(OCI)2.2Ca(OH)2, which are less soluble than ealeium hypochlorite. Sec O'Connor (l.c.), who studied the system

Ca(OCI)₂/Ca(OH)₂.

The available chlorine content of bleaching powder (35-37% commercial standard) compares very unfavourably with pure calcium

hypochlorite (99·2%).

Since 1906 the large number of patents and the very appreciable volume of work indicate the interest of the chemical industry in the production of pure calcium hypochlorite. The successful exploitation of this work is due chiefly to Chemisebe Fabrik Grieslicim-Elektron, I.G., who market a product known as "Perchloron," which contains approximately 75% available chlerine. Griesheim-Elektron, (G.P. | contains 75% or more available eblorine.

cast-iron vessels. In order to obtain complete [188524, 1906, and B.P. 8476, 1907), describe the chlorination of milk of lime nearly to saturation and concentration of the filtrate in vacuo. Crystals of calcium hypochlorite hydrato separato out, are filtered, and then dried in vacuo (see also B.P. 17859, 1914, and G.P. 282746, 1913). The product is said to contain 80-90% available chlorine and to be more stable than bleaching powder.

Mathieson Alkali Works Inc. market a product H.T.H. (High Test Hypochlorite) which contains upward of 60% available chlorine (Gage, Chem. Mot. Eng. 1929, 36, 295).
In B.P. 317233, 1928 (Mathieson Alkali Works

Inc.), it is stated that calcium hypochlorite in pure form is relatively stable, but the presence of calcium chloride in substantial amount makes it relatively unstable, owing to the hygroscopic properties of the salt.

The low solubility of basic calcium hypoehlorite may be used in the production of calcium hypochlorite to remove calcium chloride from the product at an intermediate stage:

(1) 5Ca(OH),+2Cl, $=Ca(OCI)_2 \cdot 3Ca(OH)_2 + CaCI_2 + 2H_2O$

(2) $2[Ca(OCI)_2 \cdot 3Ca(OH)_2] + 6CI_2$ =5Ca(OCI)2+3CaCI,+6H,O

The preparation of basic calcium hypochlorite is described in G.P. 195896, 1907, and 373847, 1921, B.P. 182927, 1921, and 188662, 1922, of Griesheim-Elektron; and its isolation by centrifuging is described in G.P. 451665, 1926, B.P. 276307, 1927, also of Griesheim-Elektron. A product containing approximately 40% available chlorine and 5-15% of water is obtained.

Dry basic hypochlorite has been sold under the name of "Basogrebt." The chlorination of basic calcium hypochlorite in aqueous suspension yields calcium hypochlorite which separates out under suitable conditions in crystalline form (G.P. 195896, 1907; F.P. 370863, 1906; U.S.P. 934467, 1906; and B.P. 24101, 1906). Urano and Imai (B.P. 266180, 1926) describe the production of basic calcium hypochlorite by the addition of the requisite quantity of lime to a clear solution of bleaching powder and conechtration in vacuo. In B.P. 266181, 1926, the same inventors treated the basic salt obtained according to the previous patent with water, filtered off the lime and obtained calcium hypochlorite by concentration in vacuo. The dried product contained 90-99% available oblorine.

F. N. Kitchen and I.C.I., Ltd. (B.P. 378847, 1931), and F. T. Mcchan, F. N. Kitchen, and I.C.I., Ltd. (B.P. 404627, 1932) describe cyclic processes for the production of normal calcium hypochlorito by separation of cither monobasic calcium hypochlorito or dibasic calcium hypochlorito as an intermediate stage. The mother liquor is rejected. The separated basic calcium hypochlorite is subsequently ehlorinated in the presence of additional water to produce solid normal calcium hypochlorite. The liquor separated from the crystals of normal calcium hypochlorite is recirculated to the first stage in the process, where it is mixed with additional lime slurry. The dried product

A series of patents by Mathieson Alkah | must be selected to prevent the product becom-Works Inc. describes the production of calcium ing contaminated with catalytic bodies which hypochlorite by neutralisation of hypochlorous acid with milk of lime. Hypochlorous acid is produced by chlorination of a suspension of calcium carbonate. The aqueous solution is extracted with carbon tetrachloride (containing 2% alcohol) or anyl alcohol (U.S.P. 1481039, 1924; U.S.P. 1481040, 1924; U.S.P. 1632463, 1927; B.P. 195366, 1923; B.P. 216450, 1923). The chlorine may be removed as a solid organic derivative such as acctechloroanshde, which reacts slowly with water to give hypochlorous acid (U.S.P. 1609328, 1926)

Mathieson Alkalı Works Inc, has a series of patents for the production of ealemm hypochlorite which are based on the addition of alkalı either as caustic soda er sodium hypochlorite to lime or chlorinated lime suspensions. In this way calcium chloride is replaced by sodium chloride, which results in easier removal and purification of the solid phase

$$2Ca(OH)_2+2Cl_2=Ca(OCl)_2+CaCl_2+2H_2O$$

 $CaCl_2+2NaOH=Ca(OH)_2+2NaCl$
 $CaCl_2+2NaOCl=Ca(OCl)_2+2NaCl$

The solid phase consists essentially of calcium hypochlorite hydrate and sodinm chloride, and a product of high a variable chlorine content can be obtained, which is presumably dried in succeo (G.P. 473924, 473925, 473975, 473976, 474080, 474220, 477333; B.P. 314447, 321250, 321720, 323205; 1927-1928)

An analysis of the Mathieson product is:

Availab		hlor	ine		63-0
CaCl.					0-5
Free lim	e				35
Inert co	nst	tue	nts		31.5
Water					1.5

(Gage, Chem Met. Eng. 1929, 36, 295). The production of a triple aslt,

Ca(OCI), NaOCI, NaCI, 12H,O,

by the addition of sodium chloride to a solution made by chlorination of lime is described in B P. 317437, 319727, 320182, 1927, of Mathieson Alkalı Works Inc.

The use of caustic sods to displace some of the hme used in the process of manufacture of calcium hypochlorite would appear to result in

an increased cost of manufacture, The obvious difficulties in the manufacture of

calcium hypochlorite are:

(1) The chlorination of a cream of lime. A cream of lime which is fairly fluid becomes thicker as the formation of basic calcium hypochlorite takes place until it has the consistency of porridge. Further chlorination to calcium hypochlorita causes an increase in fluidity. Great care is essential if decomposition of hypochlorite to chlorate is to be avoided.

(2) Calcium hypochlorite bydrate in the form of fine needles must be freed from a fairly concentrated solution of calcium chloride if a high-test final product is to be obtained. operation presents considerable difficulty.

would cause decomposition, e.g. salts of iron,

nickel, etc. (4) Drying of the filter cake must be earried out at low temperature, otherwise decomposition similar to that of bleaching powder on heating

will take place (Ditz, Z. angew. Chem. 1901, 14, 3, 25, 49, 105).

"Maxoelor," a calcium hypochlorite product containing 75% or more of available chlorine, is manufactured in England by Imperial Chemical Industries, Ltd. "Perchloron" is manufactured in U.S.A. by Pennsylvania Salt Co, at Wyandotte (Chem.-Ztg., Oct. 1931, 780).

Manufacture of calcium hypochlorite is earried out in Japan by Kojimachi Ku Co. (World Trade Notes on Chemicals, Feb. 1931, p. 2).

Solution Hypochlorite Potassium (Chloride of Potash or Eau de Javelle) .- This liquor was first made in 1789 at the Javel Works near Paris, and was the first bleaching compound known. It was then made by passing chlorine into a solution of potashes (crude potassium carbonate) in eight parts of water. It is no longer used, having been replaced by the cheaper sodium and calcium

hypochlorites.
Sodium Hypochlorite Solution (Eau de Labarraque, usually called Eau de Javelle) -As originally prepared, by passing chlorins into a solution of sodium carbonate untd the bound began to effervesce, sodium hypochlorite solution was very unstable. This high instability was caused by the presence of aubstantial amounts of hypochlorous acid, which is a weaker

acid than carbonic acid. Hypochlorite of soda solution is now prepared

by one of the three following processes. 1. By double decomposition of calcium hypochlorite and sodium carbonate or sulphate with precipitation of the calcium as carbonate or sulphate;

Ca(OCI),+CaCI,+2Na,CO =2CaCO+2NaOCI+2NaCI

The calcium hypochlorite may be as bleaching powder or as one of the modern " high strength calcium hypochlorites. This method is usually adopted for bleaching purposes in smaller factories.

2. By passing chlorine into a solution of caustic aoda untd nearly all the soda is chlorinated:

Substantial amounts of this solution are made at electrolytic alkali works, and sold at strengths of 10-15% available chlorine. Sodium hypochlorite may, however, be conveniently prepared sa asta from sodium hydroxide solution and

hound chlorine.

It has been found necessary to leave a small amount of free sodium hydroxide, say 0 3% in order to make the solution stable.

3 By electrolysis of hrine solution. Under Production of Chlorine, electrolytic cells devised to liberate chlorine from common salt solution have been discussed. In the case of the prepara-(3) The materials of construction of the plant tion of sodium hypochlorite there is no necessity when the electrolysis starts to keep the main i products, chlorine and caustic soda, apart; indeed, they must be brought together. This indeed, they must be brought together. This contravenes the generalisation that the products of electrolysis must be kept separate in the cell and, consequently, a sodium hypoelilorite cell compares very unfavourably with the chlorine cell with subsequent recombination of chlorine and caustic soda outside the cell.

As in chlorine cells (excluding those with mercury cathodes) the primary products of the electrolysis are hydrogen and caustic soda at the cathode, and chlorine at the anode. With the mixing of catholyte and anolyte, sodium hypo-

chlorite is formed:

$$Cl_2+2OH'=Cl'+OCl'+H_2O$$

or $Cl_2+2NaOH=NaCl+NaOCl+H_2O$.

Since sodium chloride is reformed, the net reaction

is the result of the passage of two faradays. A number of side reactions are important. Since, in spite of mixing, there will be an excess of chlorine near the anode, sodium chlorate will be formed chemically:

but, at ordinary working temperatures, this reaction is slow and unimportant. Electrolytic reactions are more important. At the eathode, hypochlorite is reduced by the nascent hydrogen:

The effects of this reaction may be lessened by the addition of potassium chromate, a vanadium salt, sodium resinate, or Turkey red oil in presence of calcium salts. These agents appear to yield a thin porous diaphragm round the enthode and so prevent the bulk of the solution from coming into contact with the active hydrogen.

If the hypochlorite ion concentration in the anode layer becomes appreciable hypochlorite ions will be released. According to Foerster and Müller (Z. Elektrochem, 1902, 8, 665), the

net reaction is:

$$6ClO'+3H_2O = 6H'+2ClO_3'+4Cl'+1\frac{1}{2}O_2+6e$$

This reaction limits the hypochlorite con-centration attainable, although loss may be minimised, at the expense of the lesser evil of chemical formation of chlorate, by not mixing efficiently in the immediate neighbourhood of the anode. Moreover, since the ratio of chloride to hypochlorito must be kept high, a large

consumption of salt is necessary.

Allmand and Ellingham ("Applied Electro-Chemistry," London, 1924, p. 375) summarise the most favourable conditions for the electrochemical production of hypochlorite solutions

(1) An electrode arrangement which will permit of an undisturbed diffusion layer being formed at the anode.

(2) A neutral electrolyte.

Vor. 111,-5

(3) Strong NaCl solution, both to lower the resistance and to permit the production of stronger hypochlorite solutions.

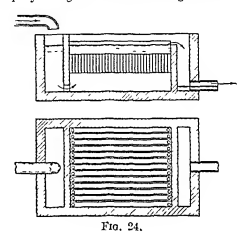
(4) Low temperature.

(5) High anodic current density.

(6) Presence of potassium chromate. (7) Use of platinised platinum electrodes.

In spite of the very formidable difficulties, several technical cells have been developed for use in districts where sodium hypochlorite or liquid chlorine is not readily available.

Kellner Cell: Vertical Type (Fig. 24).—This cell, now obsolete, consisted of a stoneware vessel divided into a number of small compartments by glass plates fitting into small grooves in its sides. The glass plates were wound with platinum-iridium wire, which formed bipolar electrodes. The electrolyte was circulated rapidly through the cell and a cooling coil.



The Haas-Oettel Cell consists of a rectangular box divided into small compartments by a series of vertical bipolar graphite electrodes set in grooves. The electrolyser stands in a tank of brine and circulation is effected automatically by the gas evolution.

The Schuekert Cell is a large stoneward trough divided into 8 to 10 compartments by means of vertical glass plates. The bipolar electrodes may be all carbon or half may be earbon and the other half platinum-iridium foil. Circulation is by means of the hydrogen evolved and is through all compartments in series, thus differing from the Kellner vertical and Haas-Octtel cells, where several passages through one compartment are obtained. Some types of Schuekert cell are provided with alternato cooling compartments.

Kellner Cell: Horizontal Type (G.P. 165486, 1902).—This cell consists of a long concreto trough divided into compartments by vertical plates of glass. The compartments are arranged as a easeade and the horizontal electrodes of platinum-iridium are bipolar, acting as eathode in one compartment and as anode in the next. The electrodes pass beneath the glass divisions and the eathode is above the anode. The electrolyte flows by gravity through all the compartments in series (Fig. 25).

based on figures given by Allmand and Ellingham (op. cit.)

G	rams avall,	kw hrs.	Tons of salt
Type	Cl _s per	per ton of	per ton of avail Cl
Kellner (vertical		6,600	12.5
Haas-Oettel .	. 123	6,500	14
Schuckert	. 20	6,100	75
Kellner (horizon	tal) 25	6,100	6

It will be seen that normally these cells cannot compete with sodium hypochlorite or liquid chlorine produced by an alkali works.

For further details, see Almand and Ellingham (op ct); or the article "Hypschlorite und elektrische Bleiche," by Bilhter and Tuchs, in Vol. 2, part 2, of Engelhardt's "Handbuch der technischen Elektrochemie"; Kurt-Arndt, "Technische Elektrochemie." Uses of Sodium Hypochlorite Solutions—

The chief use of codium hypochlorite solution

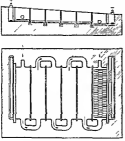


Fig. 25.

is the bleaching of high quality fabrira made from vegetable fibres. It is also used in laundries. Sodium hypochlorite solution possesses strong germicidal powers and is used very extensively as a disinfectant. It is also used

in the production of unshrinkable weetlen goods. Chlorates.—The chlorination of hot alkelt solutions gives a muxture of chlorade and chlorate in the molecular ratio of 5 to I, according to the equation:

2KOH+Cl=KCIO+KCI+H,O;

but heating the hypochlorite solution after neutralisation of the residual sikelinity rauses decomposition to chlorate. Empirically the reaction may be written

3KCIO-KCIO3+2KCI,

Comparison of Cells -The following table is but the actual mechanism is possibly very aged on figures given by Allmand and Elling complex.

MANUTACTURE OF CRIDERIUS. CRIMERIUS METHORS—Orginelly potassium chlorate was manufactured by chlorantion of potassium hydroxide solution, but this process was abandoned owing to the conversion of over the airths of the expensive potassium into relatively useless potassium chloride. A suggestion of I Lebig led to the adoption of a process depending on the chloritantion of milk of lime to estimum chlorate and calcium chloride.

$$6Ca(OH)_1+6Cl_1$$

= $3Ca(OCl)_1+3CaCl_1+6H_1O,$
 $3Ca(OCl)_2-Ca(ClO_3)_2+2CaCl_3$

followed by double decomposition of the calcium chlorate with potassium chloride to give potassium chlorate:

Since calcium chlorate, calcium chloride, and potassium chloride are all very soluble in water, and potassium chlorate is only sparingly coluble, the reaction could be carried out with lattle loss of potassium chloride.

The first stage, the preparation of calcium hypochlorite solution, was carried out in cast-iron vessels. The temperature was kept below 35°C, to decrease the decomposition of the hypochlorite to chloride and oxygen,

a reaction estalysed by iron salts.

When all the lime was neutralized sponinneous decomposition to chlorate took place with considerable rise in temperature. The crude solution of calcium chlorate and calcium chlorade was concentrated and settled or filtered. After addition of a slight excess of potasium chlorade solution, the solution was once more concentrated, settled, and the clear solution cooled. The crude potassium chlorate was pusified by recrystillustion.

Two important variations of this process to work or Nordo out. In the Pechney process to original solution of calcium chlorate and calcium chlorate and calcium chlorate and calcium chlorate and calcium chlorate had crystallized amount of the calcium chlorate had crystallized out (probably as CaClip-HQ.) In this way the amount of calcium chlorate remaining in solution could be reduced to a molecular ratio CaCli, Ca(ClO₂), of 1-2:1. There was, therefore, a greatly decreased bulk of calcium chlorate solution with consequent decreased losses of potassium sails in the mother liquors.

In the Muspratt process megnesia was used instead of lune. The molecular ratio of chloride to chlorate was only about 51:1,1,25 against 53:1 for the lune process. The luquor was evaporated and magnesium chloride heart states of the control of the

profitably.

(2) Magnesium chloride solutions are very

Sodium Chlorate.—Owing to the great solubility of sodium chlorate, manufacture by considerable eliemical methods presented difficulties.

Péchiney Process.-The chlorate-rich solution obtained in the Péchincy process for potassium chlorate was treated with solutions of sodium sulphate, precipitating all the calcium as calcium sulphate. The solution of sodium chloride and sodium chlorate was concentrated by evaporation with separation of most of the sodium chloride. The solubility of sodium chlorato increases rapidly with increasing temperature, but the solubility of sodium chloride increases little. Cooling of the chloraterich solution, therefore, yielded sodium chlorate containing relatively little sodium chloride.

Muspratt Process (see above).—The liquors, containing magnesium chlorate and reduced amounts of magnesium chloride, obtained as in the process for potassium chlorate, were treated with sodium carbonate to precipitate all tho magnesium as basic magnesium earbonate, which was used to prepare magnesia for the initial chlorination. The solution of sodium chloride and sodium chlorate was worked up as

in the Péchiney process.

The chemical methods for the manufacture of chlorates have been superseded by the electrochemical mothods.

2. Electrochemical

METHODS. — Under sodium hypochlorite (q.v.) it has been shown that chlorato may be formed electrochemically by the reaction:

6C1O'+3H2O

 $\pm 6H' + 2CIO_3' + 4CI' + 1\frac{1}{2}O_2 + 6c$ (i)

or chemically by the reaction:

$$2HClO+ClO'=ClO_3'+2Cl'+2H'$$
 (ii)

These two reactions are the basic reactions in the two types of cell which have been developed for the production of chlorates.

(i) Electrolysis of Alkaline Alkali Chloride Solutions .- Consideration of the reaction (i) above with the necessary previous reactions,

$$12Cl' = 6Cl_2 + 12c$$
.
 $6Cl_2 + 6OH' = 6ClO' + 6Cl' + 6H'$,

shows that, since hypochlorite is both formed and decomposed by electrochemical reactions, the formation of 6 gram-ions of chlorate is tho result of the passage of eighteen faradays, of which twelve are used to produce active oxygen in the form of chlorate and six yield gascous oxygen. The maximum current efficiency is, therefore, 66.7%. Moreover, since hypochlorous acid is an extremely weak acid and a maximum concentration of hypochlorite ions at the anode is desirable, the alkali chloride solution must be alkaline. If the alkalinity is excessive, trouble is experienced owing to the direct discharge of hydroxyl ions at the anode.

(1) The magnesium chloride could not be sold | divided by a porous asbestos diaphragm. cathodes were made of iron, or preferably nickel; the anodes were of platinum-iridium foil. The current density was 50 amps. per square decimeter, and the voltage 4.5 to 5.0.

Although, according to Billiter ("Techn. Elektrochemic," II, 358), the current efficiency was only of the order of 25%, this process had considerable commercial success owing to its simplicity and the availability of very cheap power.

Hurter (B.P. 15396, 1893) also described a cell for the production of potassium chlorate. by the electrolysis of alkaline potassium chloride

solutions.

These processes have now been completely processes superseded by involving electrolysis of acidic alkali chloride solutions.

(ii) Electrolysis of Acidic Alkali Chloride Solutions .- The basic reaction of the production of chlorates by the electrolysis of acid solutions of alkali chlorides is shown by equation (ii) above.

This reaction proceeds extremely slowly in alkaline solution owing to the neutralisation of the hypochlorous acid. Hypochlorous acid, is, bowever, a very weak acid, and the formation of chlorato may be considerably accelerated by making the solution weakly acidic. Moreover, the temperature coefficient of the velocity of reaction is high, so that increase in temperature causes a very great increase in the rate of formation of chlorate. Since formation of chlorate by the discharge of hypochlorite ions is disadvantageous (wasting energy), it is desirable to have a largo volume of electrolyte compared with cleetrode area or total current, as the chemical formation of chlorate proceeds throughout the electrolyte, and is not restricted to the neighbourhood of the electrodes. Cathodic reduction is best avoided by the addition of a small amount of potassium chromate.

Originally platinum or platinum alloys were extensively used as anodes. Graphite is rapidly attacked at the high temperatures (80°-90°C.) originally employed, but the very great fluctuations in the price of platinum have led to the use of graphite at relatively low temperatures (40°C.).

1. Barker Cell.-In the cell developed by J. T. Barker for the United Alkali Co., the container is a narrow iron trough, the lower part being covered with cement. The cathode consists of the exposed part of the iron trough and a central iron cooling coil. The anodes are vertical, cylindrical graphite rods. To decrease oxidation of the anodes, the graphite is impregnated with paraffin wax or other inert organic material. The cell top is a plate of an asbestos composition and the anodes are supported by ebonite rods passing through the tops of the graphite and resting across the top of the trough.

A 1,500-amp. cell is about 8 ft. long, 4 ft.

deep, and 8 in. wide. The working temperature is about 40°C (Fig. 26).

2. Aussiger Verein Cell.—In this cell special attention has been paid to the use of a large volume of electrolyte for a given electrode area, Gall and de Montlaur (B.P. 4686, 1887) so that the chemical production of chlorate may described a process for the production of be accelerated while electrolytic discharge of alkali chlorates. A stoneware trough was

quarter of the concrete trough is occupied by the lat the top of the electrode than at the bottom, electrodes which consist of alternate graphite This gives more even wear and longer life of the slab anodes and iron stim eathbods. All the anodes. anodes are connected to a graphite block, from which a graphite rod rises to above the electro- covered iron. The temperature is maintained The eathode connection is an iron bar at 40°-50°C, by circulation of the electrolyte, passing through the side of the cell to which the individual eathodes are fastened.

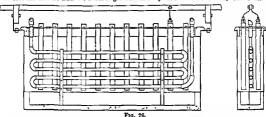
3. Angel Cell -In this cell also the volume of electrolyte is large, but the electrodes are suspended at the top of the cell with the space for the chemical reaction at the bottom Graphite anodes and iron cathodes are used, and a special crystallising or grinding chlorates. The use of feature of the cell is the setting of the cathodea wooden vessels for erystallising potassium relevant to the anodes If the electrodes are chlorate has in the past led to disastrous fires set parallel, their resistance causes most of the and explosions. current to pass between the tops of the electrodes, with consequent rapid destruction of the in the manufacture of matches and explosives tops of the graphite anodes. In the Angel cell, and in the development of dyes. A recently therefore, the iron cathodes are set at an angle developed use is that of sodium chlorate as a so that the distance from the anode is greater relatively non-toxic weed killer, but a dis-

The cell container is either iron or lead-

Safety Precautions in the Manufacture and Use of Chlorates -Owing to the very high proportion of available oxygen, chlorates are dangerous in contact with combustible materials. In Great Britain special washed elothing must

he provided for every worker engaged in

Uses of Chlorates - Chlorates have been used



advantage is the risk of fire when wood or other mibstances

Perchiorates. - Although potassum and important than the hygroscopic sodium per-chlorate, sodium chlorate is the usual raw material since its great solubility is advantageous for the electrolytic preparation of sodium per-Although the empirical sum of the chlorate cell reactions may be written

the actual mechanism of the reactions is very complex, chlorites being probably formed as an intermediate, while chlorides are inevitable by products of the reactions. The mechanism of this formation has been represented as the result of.

(1) cathodic reduction of chlorate, (ii) chemical decomposition of chlorate, 4NaClO,=3NaClO,+NaCl,

The second of these possibilities is probably quite unimportant.

As electrolyte a concentrated solution (60comhustible material is left in contact with these 70%) of sodium chlorate is used. The cathodes are iron and the anodes amouth platinum. A very high anode potential is essential and, ammonium perchlorates are technically more therefore, graphite and magnetite are unsuitable. Normal cathode current densities of 10-20 amps. per square decimetre are used, but the anode current densities are extremely high, ranging from 30 to 70 amps per square decimetre. Originally low temperatures were used, but modern practice allows temperatures as high as 60°C.

The discharge of perchlorate ions, with consequent release of oxygen and drop in current efficiency, is not serious until very high perchlorate chlorate ion ratios are reached. fact is important, since potassium perchlorate and potassium ehlorate form mixed erystals, and a very high sodium perchlorate sodium chlorate ratio must, therefore, be obtained in the electrolytic cell.

In practice, current efficiencies are about 80-

Perchloratea are used extensively in some types of explosives. In recent years perchloric acid has been used as an important analytical reagent (c. CHEMICAL ANALYSIS, Vol. 11, p. 584d), and magnesium perchlorate) anhydrone) as a drying agent in analysis.

Bibliography.—Allmand and Ellingham, "Applied Electrochemistry," 1924; "Handbuch der technischen Elektrochemie," vol. 2, part 2, edited by Engelhardt, article "Die elektrolytische Perchloraterzeugung," by Billiter and Fuchs; Winteler, Z. Elektrochem. 1898, 5, 50, 217; 1901, 7, 635; Ocensli, ibid. 1903, 9, 807; Blan and Weingand, ibid. 1921, 27, 1; Coulern, Chem.-Ztg. 1906, 30, 213; Bennett and Mack, Trans. Electrochem. Soc. 1916, 29, 323; Howard, ibid. 1923, 43, 51; Williams, Trans. Faraday Soc. 1920, 15, 134; Knibbs and Palfreeman, ibid. 1921, 16, 402, 424; Foerster, "Elektrochemic wässeriger Lösungen."

Hydrochloric Acid,—Commercial hydrochlorie acid is a more or less pure solution of

hydrogen chloride, HCl, in water.

Hydrogen Chloride is a colourless gas, of sp.gr. 1.2596 (air=1) and molecular weight 36.464. Its ni.p. is -111.4°C, and b.p. -85.0°C, and its critical constants are: temperature, 51.5°C.; pressure, 81.6 atmospheres; density, 0.424. Vapour pressures greater than one atmosphere are given by the formula

$$\log_{10} p = 7.09 - \frac{1055}{T} - 0.0105T + 0.000014T^{2}$$

where T is the absolute temperature.

The heat of formation of a grain-molecule of

hydrogen chloride is 22,000 cals.

Hydrogen chloride is very soluble in water, the solution being commonly known as hydrochloric acid, muriatic acid, or spirits of salt.

The following tables are abstracted from "International Critical Tables," to which reference should be made for more detailed statements of physical properties.

Weight Percen- tage HCI.		Density (g, per nili	illitre) at	
her tage	orc.	10°C.	20°C.	50°C.	100°C.
1 2 4 6 8 10 12 14 16 18 20 22 24 26 30 32 34	1·0052 1·0106 1·0213 1·0319 1·0423 1·0528 1·0634 1·0741 1·0840 1·0958 1·1067 1·1177 1·1287 1·1396 1·1505 1·1613	1·0048 1·0100 1·0202 1·0303 1·0504 1·0507 1·0711 1·0812 1·0920 1·1025 1·1131 1·1238 1·1344 1·1449 1·1553	1·0032 1·0082 1·0181 1·0279 1·0376 1·0474 1·0574 1·0576 1·0878 1·0980 1·1083 1·1187 1·1290 1·1493 1·1493 1·1593 1·1593	0·9929 0·9977 1·0073 1·0168 1·0262 1·0357 1·0453 1·0549 1·0646 1·0743 1·0937 1·1033 1·1129 1·1224 1·1318	0.9636 0.9688 0.9791 0.9892 1.0090 1.0188 1.0286 1.0383 1.0479 1.0574 1.0668 1.0761 1.0853 1.0942 1.1030
36 38 40			1-1789 1-1885 1-1980		

For density determinations by hydrometers, see B.S.S. 718, 1936.

VAPOUR PRESSURES OF HYDROGEN CHLORIDE SOLUTIONS AT 25°C.

Weight percent- age of HCl.	Partial pressure of water in nim. Hg.	Partial pressure o HCl lu nim. Hg.
6	21.8	0.00131
10	20.0	0.0067
14	18.0	0.0316
18	15.4	0.148
20	14-1	0.32
22	12.6	0.68
$\frac{22}{24}$	11.4	1.49
26	9.95	3.20
28	8.75	7.05
30	7.52	15.1
32	6.37	32.5
34	5.35	68.5
36	4.41	142
38	3.60	277
40	2.88	515

The behaviour of aqueous solutions of HCl under varying conditions of temperature and pressure has been studied by Roscoe and Dittmar (J.C.S. 1860, 12, 128). They found that concentrated hydroehlorie acid on heating loses both gas and water, but as the proportion of HCl to H₂O is higher in the vapour than in the original liquid, the residual liquid gradually becomes weaker, until a sp.gr. of 1·101 and a percentage of 20·24 HCl are attained at a boiling temperature of 110°; the acid in this state distils without change, provided the atmospheric pressure is 760 mm. At a lower pressure the acid distilling unchanged is stronger, at a higher pressure it is weaker. A similar, but not identical, limit is attained when strong acid is exposed to the air at ordinary temperatures.

The table (p. 70a) from Lunge and Hurter, "Alkali-Maker's Pocket Book," 1884, gives the percentages of HCl in aqueous hydrochloric acid of various specific gravities and at a

temperature of 15°.

Lunge and Hurter's "Alkali-Maker's Handbook" 1891 gives somewhat lower figures. The table in the right-hand column of p. 70 gives corrections of the hydrometer readings for

varying temperatures.

The Manufacture of Hydrochloric Acid.—The Leblanc soda process, which had been gradually diminishing in importance since 1902 in Great Britain, ceased to be operated in 1920, although considerable quantities of saltcake are still manufactured for the sulphate pulp, glass making, and sodium sulphide processes (see "Merseyside," S.C.I., Liverpool, 1924). This decline of the Leblane process provided a growing output for hydrochloric acid produced from electrolytic chlorine and hydrogen. There was a rapid increase in the manufacture of hydrochloric acid produced by combustion of chlorine in hydrogen which had been already commenced by Castner-Kellner Alkali Co., Ltd., in 1912. This combustion on the principle of the Bunsen burner may be carried out in open or closed burners with an excess of hydrogen.

Deg	Spec	Per-	Grams	foot	foot con-	a l
dell.	gravity.	of HCI	per litre	weighs	tains lb.	Ι,
		0	Jan Miles	Ib.	of HCI.	1
1	1.005	1.12	11 32	62 66	0.70	i i
2	1-010	2 12	21-45	62 97	134	
3	1 015	3 12	31 67	63 29	1 97	
4	1 020	411	4199	63 60	261	li
5	1 025	5-11	52-41	63 91	3 26	l i
В	1 030	6 11	62 93	61 22	3 92	Ι.
7	1-035	7 10	73 55	64 53	4 58	
8	1 040	8 10	84 27	64 84	5 25	Ι.
9	1.045	9 10	05 09	65 16	6 93	1
10	1 050	10-09	106 01	65 47	661	
11	1 055	11 09	117 02	65 48	7 31	ı
12	1 060	12 09	128-14	66 09	8 00	ı
13	1 065	13 09	139 36	66 40	8 69	ير ا
14	1 070	14 08	150 68	66 71	9 40	₽
15	1 075	15 08	162 10	67 03	10 11	A G
16	1 050	16 07	173 62	67 34	10 83	
17	1 085	17 07	185 24	67 65	11 55	ΙĔ
18	1 090	18 07	196 96	67 96	12 28	۱ă
19	1 095	19 07	208 76		13 02	15
20	1 100	20 06	220 70	68 59	13 76	12
21	1 105	21 06	232 68	68 90	14 51	lê.
22	1 110	22 06	244 80	69 21	15 27	Пуввоситовія
23	1 115	23 05	257 02	69 52	16 04	17
24	1 120	24 05	269 34	69 83	16 79	8
23	1 125	25 05	281 76	70 14	17 57	. ج ا
26	1 130	26 04	294 26	70 46	16 35	GRAVITY
27	1 135	27 04	306 90		19 14	13
29	1 140	25 04	319 62	71 03	19 93	I۵
29	1 145		332 44	71 39	20 73	
30	1 150	30 03	345 36	71 70	21 54	Sprente
31	1 155	31 03	358 34	72 02	22 35	Į₽
32	1 160	32 02	371 44	72 33	23 16	١٤
33	1 165	33 02	384 64	72 04	23 99	l"
34	1 170	34 02	397 94	72 95	24 82	Į,
35	1 173	35 01	411 34	73 26	25 65	
39	1 180	36 01	424.84		26 49	ē
37	1 185	37 01	436 44	73 69	27 34	
38	1 190	38 01	452 14	74 20	28 20	Ĕ.
39	1 195	39 00	466 00	74 51	29 66	ATURE
40	1 200	40 00	479 84	74 82	30 00	ě
			1			PEWFER
						12
lu the	obeu pr	irner exc	ess hydr	es ango	burnt by	
dmis	to nor	air ab	ose the	cplorm	s flame,	5
reult	mg in a	n absor	ption sy	stem fr	eo from	
the d	mer of	explosio	n Int	ne closed	1 burner	FLUENCE
ercess	hydrog	en carn	es the h	y drochk	rie acid	Ě,
KAS TO) the ad	*crption	system.	Great	t care is	15'
uc res	ary in th	us case t	o avoid e	Thiosion	2010	
HORS	or sydre	emorie i	sero men	e irom	gas thus	-
bre.In	red mas	with care	be produ	aced of F	naly tical	١.

reagent quality In former days small quantities of acid were

produced by decomposing common salt with sulphirm and in cast iron cylinders, and in order to utilise completely the relatively expen sive sulphurk acid an excess of common salt was used. The solul product left in the cylinder consisted of sodium sulphate mixed with 20-30% NaCl and was commercially known as cylinder cake This method of production of hydrochloric acid has, however, practically ceased

Some time after the establishment of the father a sla process in this country all the

Per- Grams 1 cubic 1 cubic bydrochloric acid of commerce was obtained as

	D. L	rout	ict	OI	I.	na	t	pr	DC	253	,	11	ПC	h	at	firs
	901	1.12	25	1 088	1 070	1.070	1 061	1 053	0.55	1 037	1 025	1013	100	0 991	0 081	0 971
	•20	1 123	125	1 090	1 081	1 073	1 064	0.25	1 047	1 033	1 027	1 015	1 003	0 993	0 983	0 973
	•00	1.125	10	1 093	1 081	1075	1 086	1 057	1 049	101	1 029	1 017	1 002	0 992	0 082	0 975
	2	1 127	28	1 093	1 080	1 078	1 063	1 079	102	1043	1 031	1 019	1 007	0 997	0 987	0 977
é	80	8	100	1 097	1 089	1 080	1 07	1 061	1 053	1 044	1 033	3	-000-I	0 999	0 080	0 979
RIG AC	22	=	: :	9	1 091	1 083	1 073	1 063	107	1 048	1 03	1 033	<u>=</u>	1001	0 091	0 081
ON THE SPECIFIC GRAVITY OF HYDROCHLORIG ACID.	<u>6</u>	25	32	100	1001	1 085	1 076	1 065	1 056	1 018	1 636	1 025	101	188	0 993	0 983
or llyn	82	213	25	9	1 096	1 088	1 073	1 067	1038	1 049	1 038	1 027	1 016	100	0.595	0 982
AVITY	ş	821	22	101	1 093	980	960	1 069	8	3	1 040	1 039	1 018	1 008	0 998	0 988
ric Gr	\$2	12:	2.2	9	5	1 093	1 082	1 62	1 063	1 053	3 642	1 03	1 050	100	1 000	06 0
Srrci	20	=	- 1	=======================================	1 103	100	1084	1 073	1901	1 075	101	1033	1 022	5	1 30	0 992
0 T	ş	=	5	=	7 100	1 097	7 088	202	990	1 057	3 016	2032	1 025	101	200	0 991
ATURE	ģ	11.147		-	108	6601	ä	Ē	_	_	Ξ	Ξ	_	Ξ	1 007	•
TEVER	35°	1.149	133	1:13	=	101	900	1 080	1 070	00	1 020	1 050	1 030	1019	000	666.0
b	30	1163	222	1:132	=======================================	1 103	1 093	1 082	1 073	1 063	1 053	1043	1 035	1 022	3	9
NFLUENCE	220	1 154	135	1 125	1112	1.103	1 095	1 081	1 075	1 006	1 055	501	103	1 024	10	00
-	20	1.157	35	1.127	112	1.107	1 097	1 087	1 077	1 067	1 057	1 047	1 037	1 0 27	1017	1 007
	ā	1 100	1.0	1.130	1:120	=======================================	130	1 000	1 080	1 050	1 000	1 650	2	1 030	20	1 010
	è	1.163	35	1.133	=======================================	1.113	1.103	1 093	1 043	1 073	1 063	1 053	1 0 23	1 0.33	10,3	1 013
	2	1.165	153	÷	÷	=======================================	Ξ	_	1 043	_	1 061	1 055	1055	=	-	1 012
	å	1.168	1.158	1.139	1.128	1.18	1.104	1 038	1 088	1 078	1 068	1 058	1 048	1 038	0.3	810

produced hydrochloric acid gas in great excess; of that which could be sold as hydrochloric acid liquor. The difficulty of the situation was aggravated by new sources of hydrochloric acid which arose when acetio acid began to be chlorinated in great quantities by the Badische Anilin und Soda Fabrik for the manufacture of monochloracctie acid and artificial indigo, and one organic compound after another came to be chlorinated. Lastly, since the output of chlorine from the electrolysis of the chlorides of potassium and sodium was not sufficiently readily taken up, processes converting electrolytic chlorine and hydrogen into hydrochloric acid are now

operated on a very large scale.

Hydrochloric Acid obtained in manufacturing Sodium Sulphate (Salt Cake) .- When Leblanc decomposed salt in order to obtain sodium sulphate he proposed to convert the hydro-chloric acid gas evolved into ammonium chloride; but when the Leblanc soda process was introduced into this country that part of his suggestions was not earried out, and the acid gas was regarded as possessing no value, and it was allowed to escape into the air. This continued to be the ease even after that manufacture had attained a comparatively large extension, but the damage caused to the neigh-bourhood by the acid gases soon brought about the intervention of local authorities and courts of law, and manufacturers had to pay com-pensation for the damages they had done. Repeated penalties compelled the alkali manufacturers to scok means for condensing the acid gas. This was first successfully done in 1836 by Gossage's coke towers, but for many years after this there was still considerable damage done by escaping acid gas. The continued disregard by the old alkali manufacturers for the rights of their neighbours led in Belgium, in 1856, to a Parliamentary Commission, and in this country in 1863 to Lord Derby's Alkali Act, and the several subsequent Acts, by means of which British alkali makers were compelled to adopt efficient means for condensing their hydro-chloric acid, and similar legislation has been to the benefit of other manufacturing countries.

The "Annual Report on Alkali, etc., Works" made by the inspectors under the Alkali Acts forms an invaluable record of all the various steps taken for improving the condensation of hydrochlorie acid, and have helped to spread a better knowledge of that process and have caused manufacturers to improve it up to the highest

standard of efficiency

The Alkali Act of 1863 made it incumbent on manufacturers of salt-cake to allow no more than 5% of the hydrochlorie acid to escape into the atmosphere. An additional Act of 1874 prescribed that no more than 0.2 grain of HCl might be present in a cubic foot of the gas escaping from the factory into the air. This corresponds to 0.454 g. HCl per cubic metre, or about three ten-thousandths by volume. The latter clause applies also to the chimney gases and not merely to the gases escaping directly from the condensing apparatus. Although these requirements were at first

they were quite possible, and the escapes registered by the Alkali Inspectors have averaged far below the limits assigned by the Acts. In 1885 only 2.13% of the total hydrochloric acid was allowed to escape, and the chimney and other gases escaping into the air contained only 0.10 grain HCl per cubic foot. No change has been made in the limit fixed in 1874, and the working of that Act remains satisfactory. Compare the survey of these Acts by the Chief Inspector under these Acts in J.S.C.I. 1892, 11, 120.

Although the various processes and apparatus used for the manufacture of sulphate of soda produce gases differing within wide limits in temperature and concentration of HCI, an efficient condensation, i.e. practically consulte absorption of the HCl in water to form a liquid acid of 1.15-1.17 sp.gr., is usually

obtained. The efficiency of the condensation of gaseous HCI in water is dependent on three considera-

1. Temperature of the gas and of the liquid. 2. Concentration of HCl in the gas s to be serubbed and of the acid liquid to be readuced.

3. Time of contact between the gas and the liquid, and the average distance between the

gaseous particles and the liquid particles, and also the thickness of the liquid layer, and on its motions internal and as a

On issuing from the sulphate of soda plant the acid gases require cooling. The apparatus used for this purpose depends on the temperature of the

gases.

If the gases issue from the pot or pan of a handworked salt-cake furnace, the temperature is comparatively low. In this ease the cooling is usually done in a row or rows of carthenware pipes. The pipes used are often unglazed, but are rendered impervious to moisture and gas by boiling in tar. In some cases the earthenware pipes are substituted by glass pipes of 12 in. diameter, made stightly taper, as shown in Fig. 27. The glass pipes arc, of course, more liable to breakages by accident, and also through the effect of violent changes of temperature, than the earthenware pipes, but in sheltered positions they last



Fig. 27.

For the hotter gases issuing from an ordinary muffle-worked sulphate of soda furnace or from the cylinders of the Hargreaves process cast-iron pipes are commonly used to cool the gases to a safe temperature before passing them as before into earthenware or glass pipes. The action of the hydrochloric acid gases on east declared by many to be impossible of fulfil- iron is very sight at a temperature well above ment, the actual results proved at once that the condensing point of liquid acid.

When open furnaces, whether "hand" or purpose of collecting any highld deposited, "mechanical" (such as the Mactear furnace), are The cisterns are usually made in this country

used, the acid gases are mixed with the products of combustion, and are consequently much hotter and more dilute. Under these conditions extensive series of east-iron pipes are used for the preliminary cooling before passing the gases into the earthenware pipes for final cooling

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The earthenware pipes used are usually 12 in. or 15 m. diameter socket pipes, the joints being made with a cement of tar and China clay. The cast iron pipes vary from 12 to 20 in in diameter, according to the quantity of gas and the arrangement of the cooling system of pipes,

they are socket pipes, the joints being made by ramming into the socket iron filings moistened of educious sandstone found in *forkshire, and with ammonium chloride solution known as "Yorkshire flag." The eisterns are

The arrangement of the cooling pipes depends on the relative positions of the salt cake furnaces and of the and condensers and the space nvailable When the salt cake furnace is a long

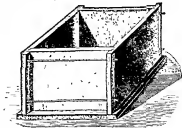


28. I'ıc

distance from the condensers, a single direct line of pipes may be a sufficient means of cooling. where the distance is small the necessary amount of cooling surface is obtained by adding



down and that start from a small stone eistern water is allowed to trickle, and up which the



composed of flags

4-6 in thick, bound together by strong iron ties, and the joints made tight by thick indiarubber cord laid in a groove between the faces of the two atones or by a mixture of tar and fireclay, as in Γigs 28, 29, 30 By one or more of the above arrange ments of pipes and custerns the acid gases are cooled to 45°, or even to 25° The cooling of weak gas is far more im portant than the

cooling of atrong gas The gases next enter the apparatus where the bydro chloric acid gas is to be converted into a hquid acid of commercial atrength. and that in such a manner that practi cally hardly a trace of acid escapes con-

densation The apparatus almost nniversally used are tall stone towers, packed with broken coke

110. 31

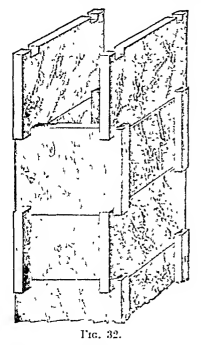
extra lengths of pipe that run vertically up and specially arranged, down which a stream of and return to another atone cistern for the gases are caused to flow (Fig. 31). These

in 1836.

The coke tower has the merits of simplicity and, when carefully packed, of efficiency.

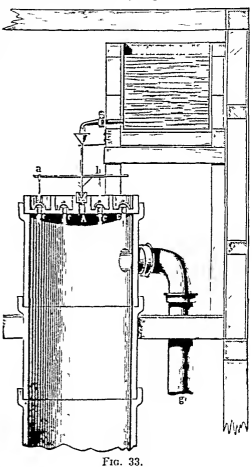
The comparative efficiency of various modes of treating liquids with gases has been dealt with in a series of papers by Hurter, who also gives some interesting data on coke-packed towers (J.S.C.I. 1885, 4, 639; 1887, 6, 707; 1893, 12. 227, 989. See also ibid. 1919, 38, 75T; 1920, 39, 23; Ind. Eng. Chem. 1924, 16, 1215; Chem. Met. Eng. 1923, 29, 146; Trans. Amer. Inst. Min. Met. Eng. 1919, 12, 231; 1929, 22, 165).

Coke towers are erections of considerable height, varying from about 40 ft. to 60 ft. horizontal sectional area may vary from 25 to 180 sq. ft. They are provided with a grating a few feet over the bottom, on which rests the coke which forms the "packing" of the



tower. This coke, which must be of as hard a quality as possible (only the best oven-coke is applicable for this purpose), is disposed in such a way that the gas is as much as possible divided in a large number of channels (none of them being so wide that the bulk of the gas cin pass up through it), and that the current of gas is constantly changed in direction. For this object long pieces of coke are laid parallel in a row; the next row is made to cross the lower one, and so forth. The size of the pieces is gradually diminished from bottom to top. In order to avoid an excessive quantity of the gas passing up the side of the condenser along the wall, it is usual to place a layer of finer coke round the condenser for a .ew inches from the wall. By this arrangement there is effected

towers were invented and patented by Gossage, | spreading of the feeding water over a very large surface. One great advantage coke has over flints or other packings is that, owing to the roughness of its surface and its porosity, it always retains a large store of liquid, which tends to keep the working of the condensers steady under a fluctuating flow of acid gases, and to tide over difficulties occasioned by any unintentional stoppage of the liquor supply. The water (or sometimes weak acid) is fed in at the top by some contrivance assuring a uniform distribution over the whole area of the tower; and in trickling down it keeps the surfaces of the coke moist, and a very large area of contact



between the gases and the liquid is thus produced. These towers are so simple to construct and so effective that of recent years there has been a growing tendency in England to do away with all cooling pipes, eisterns, wash towers, etc., and to use a couple of very large stone towers to do the whole work of cooling and condensation for two or even more salt-cake pots and furnaces. The towers are worked in series, the weak acid from the second or final tower being run down the first tower. In this way both complete absorption of the acid and the production of a both a great division and constant mixture of strong hand acid, i.e. one of over 1.15 sp gr., the gases, and at the same time a suitable are assured. To condense the acid from two

hand salt-cake furnaces the two towers may foundation, even if it amounts to much less each be 5 ft. aquare and 50 ft. high. No difficulty is now experienced in pumping

the weak acid made in the second condenser on to the top of the first condenser, either by stoneware ram pumps, ebonite ram pumps, stoneware acid eggs, or occasionally even yet hy the Harlehurst membrene pump, patented in 1876. The ascending pipe is usually of atoneware or ebonite, occasionally of lead.

Brick towers are not to be recommended, as it is practically impossible to keep the numerous joints so tight that no seid oozes out. Columns of stoneware pipes cannot be made of a large area, and, therefore, are not very well adapted to large works, at least if intended for coke towers; hut they do very good service at smaller works, or as aupplementary to a large condensing plant of stoneware receivers. The material mostly employed for coke towers is send-proof sandstone or, in France, Volvie lava, just as



for acid tanks. The system of joining the flags together is the same as shown in Figs. 28 and 29, or else as in Fig 30, or as shown in Fig. 32, in which the flags are kept together by means of grooves worked in the stone, and without the aid of any ironwork, which is always a source of anxiety in acid condensers.

The coke towers are generally placed at a high level, and are, if neressary, put upon pillars, for the purpose of running the acid from them by gravitation either into chlorine stills or into acid wheels and have introduced troughs prostore cisterns for sale. They must, of course, have a very secure foundation, in the construction of which not merely the great weight of the lutes. This arrangement is indicated in Fig. 31. tower must be considered but also the proba. This figure represents a tower hull as Fig. 32, tower must be considered but also the proba-hility that there will be now and then leakages that the considered many make the foundation unsafe. This can be avoided by esphalting the ground in such a say that any acid running down in such a say that any acid running down. cannot penetrate into the soil, but is conveyed plate tower, shown in Figs. 35, 36, and 37. to a safe distance. Any damage done to the lt consists of an earthenware tower packed

altogether, is extremely injurious, because it will almost unavoidably lead to the tower getting out of level, in which case the hquid will principally descend on the lower side; thus the necessary distribution of the liquid is not attained, and the action of the condenser is most injuriously affected.

It has been already stated that the feeding of the towers with water or with weak acid

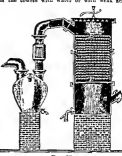


Fig. 35.

must take place in such manner as to distribute it equally over the whole surface of the coke Formerly this was generally done by a distri-buting wheel (shown in Figs, 33 and 34) worked by the pressure of the feed water itself. Fig. 33 shows two retating jets (a and b), feeding the outer and mner circles of holes, causing the stream of water to reach successively every one of the holes on the top of the condener,



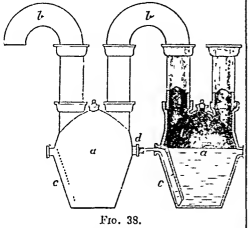
Fro. 36.



and thus securing a uniform distribution independently of an exact level of the top of the tower. Most manufacturers have discarded these uded with a number of overflows all at the same level, feeding the same number of water-

75 CHLORINE.

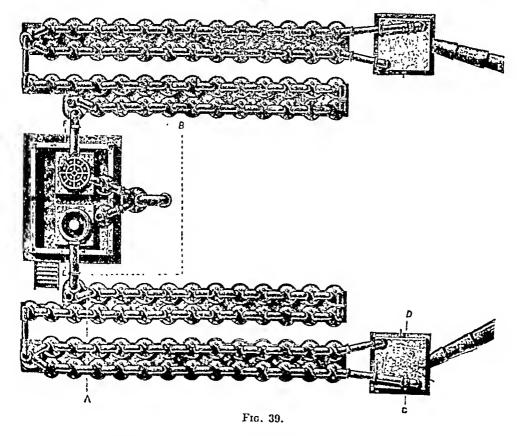
forations being made to alternate in successive plates, so that each hole corresponds to a solid



place in the plates just above and below. Figs. 36 and 37 show one of the plates in detail.

with a series of perforated plates E, E, the per- against the improved packings for the condensing towers is very conflicting, and as the cost of them is naturally higher, the coke-packed tower has fully maintained its position.

In France and Germany a somewhat different system of eooling and condensation is in general use. The earthenware and glass pipes mentioned above as being used almost exclusively in this country for the cooling of the gases are more or less completely substituted by a series of earthenware Woulfe's bottles, known technically as bombonnes or tourils (see Fig. 38). The scale of the drawing is too small to show that the vessel on the right is about 1 in. higher than the vessel on the left; the liquids travel downhill from right to left, and consequently tho gases are made to travel from left to right. The size of the bombonnes varies from 40 to 100 gallons. They are arranged in sets of 40 up to 100, as shown in Fig. 39, and, in spite of the relatively high temperature in the leading bombonne, eventually give a liquid acid of 1·15-1·18 sp.gr. They are made of the best stoneware, which is able to stand somewhat violent changes of temperature. After leaving the bombonnes, the acid gases are finally serubbed with water in a coke-packed tower, As is to be expected, the evidence for and similar in design but usually much smaller than



those used in this country. The weak liquid figure does not show the means of running off acid produced in this tower supplies the stream the strong acid from the lowest bombonnes in of weak acid required by the bombonnes. The the series, nor the cisterns, etc., for receiving the

seed. Coolers and absorption plant can now references given for the production of chlorine be made of fused silica (Vitreosil) in the form of from these bodies earlier in the article. So fer

of plant required is effected

In the condensation of hydrochlone acid gas in water with the apparatus at present used and described above, there is little reason for uspours into the neutral salt of commerce. lifficulty in avoiding losses of HCl through in-complete absorption. The difficulty more rommonly experienced is to combine the complete theorption enforced by law with a high strength iquid acid.

The impurities found in commercial hydrothloric acid made by the above processes are, hiefly, sulphuric acid, ferric chloride, arsenie nebloride, and either free chlorine or sul

shurous acid

Sulphuric acid is the largest impurity. The ly drochloric acid condensed from the pan or pot tas is always much purer than that from the paster gas. The quantity of sulphuric acid in he gases from the salt cake furnaco sometimes imounts to 2% and upwards, it was very roublesome in the Weldon, and fatal in the Deacon, chlorino process, and in some other patents taken out in 1886) applications of hydrochloric acid. The sul phuric acid in the salt cake furnace gases can be removed to a large extent by a small conlenser or dry scrubber before condensation. A large number of suggestions has been made or the purification of the salt cake furnace cases, so as to render them suitable for the Deacon process, but the only one of these proresses to achievo practical success is the Hasen elever process (BP 3393, 1883) (See at end

of Deacon Chlorine Process) Arsenie gets into the hydrochlorie acid through the sulphuric acid employed in its manufacture, the arsenic is thereby converted into AsCl₃, and passes as such into the conden-sing apparatus. Nany plans have been proposed for its removal-most frequently a treat ment with sulphuretted hydrogen or with sulphides Bettendorff (Dingl poly J 1569, 194, 253) precipitates the arsenic from concentrated hydrochloric acid by stannous chloride, and then distils the decented acid. Duflos dilutes the acid to sp gr 1 13, and digests it with strips of sheet copper at 30°C for 24 hours, and repeats this treatment with freshly scoured copper This removes all the arsenie and the free chlorine, and reduces the ferric chloride to ferrous chloride, which remains behind on distillation Beckurts (Fischer's Jahresb 1884, 30, 348) distils hydrochloric acid with ferrous chloride and removes the first 30% containing all the arsenic.

Selenium is sometimes found in such quan tities in hydrochloric acid as to give trouble

(1) from CaCl2, (11) from MgCl2; (111) from NH C! In this connection compare the The applications of hydrochloric acid include

seemade of insect since (vitered in these can be as is known, none of the processes is used, water couled by sprays a reduction in the size Mond (1883) heated the ammonium chloride with so much sulphuric acid that the acid sulphate was formed and all HCl expelled. The acid sulphate was converted by means of ammoniacal O. N. Witt (1886) expelled the HCl from NH,CI by means of syrupy Phosphoric acid. which formed ammonium phosphate. On hesting this at a higher temperature the smmonia was driven out and the phosphoric acid re-generated Jurisch (Dingl. poly. J. 1888, 267, 431) has shown that this process is not prac ticable, because there is no material known which resists the molten phosphoric acid, and only 63% to 86% of the NH₃ is recovered. Mond passed the vapour of ammonium chlorade over nickel oxide, heated to at least 350°, which absorbed the HCI and allowed the NHa to pass on. After a certain time the process was changed by raising the temperature to 500° or 600° and decomposing the nickel chloride by steam, when HCl was driven off and NiO regenerated.

> During recent years, when the chlorination of organic compounds has advanced by vast strides, the hydrochloric acid gas produced when a hydrogen atom is substituted by a chlorine atom has to be collected and utilised, The hydrochloric acid gas evolved is almost pure, easily condensed, and yields an acid quita free from the usual impurities

Hydrochlone and has also been produced by passing chlorine over wood at a temperature of 150°-350°, whereby chercoal is also formed (Bosman Electricity Works at Jaica, GP. 158086)

Lastly, hydrochloric acid has been prepared by processes inverting the practices of the last hundred years, namely, it has been prepared from its elements as a direct and definite pro cess without the formation of Ly-products, or for any other reasons than its own preparation and the utilisation of the two elements formed by the electrolysis of potassium and sodium chlordes The gases are mixed and passed over charcoal (G.P. 114219), or they sro burnt together at jets (Soc. Itahana di Elettro-chimica, JSCI 1915, 34, 1145), or they are combined together by Roberts' Patent (Electro-chem and Met Ind 1910, 8, 701), and by this last process the acid has been manufactured in

Nagara sinco 1914. The conveyance and storage of hydrochlone acid of atrengths varying from 27 to 40% has undergone changes in recent times following improvements in the technique of the appliestion of rubber preparations to metal and other in its application (Davis, J.S.C.I. 1883, 2, surfaces, Steet tanks inted with material sarying 157).

Hydrochloric Acid from Sources other than the for conveyance of send in bills (Chem, and Ind. Lellane Process—Attempts have been made to 1924, 43, 1804; 1932, 51, 205, 224; Chem. Produce hydrochloric acid from many sources, Met Eng. 1923, 35, 673; 1929, 36, 152; Ind. notably from the residual liquors of the ammonia. Eng. Chem. 1927, 19, 130; Ind. Chem. 1928, 4, soda process. A number of suggestions has 75) The familiar glass carboy is still, however, been made for the direct production of HCI: used to a large extent for the conteyance of this acid.

by neutralisation in solution and subsequent tion of aldehyde, as follows: crystallisation, and by the dry process involving sublimation described in B.P. 273093, 274263, 290045 (Moore, Polack and Castner-Kellner Alkali Co., Ltd.). The older application of hydrochlorie acid for the production of chlorine by the Weldon and Deacon processes is no longer of importance in Great Britain.

It is used for extracting the phosphates from bones in the production of osseine for gelatine; for preparing zinc chloride, stannous chloride. and ferrous chloride; for the preparation of pure carbonic acid from limestone; for the manufaeture of aniline hydrochloride; and in the manufacture of many coal tar colours and intermediates.

In recent years the operation of eleaning (pickling) metal with hydrochloric acid prior to rolling into sheet, wire drawing, and coating with zinc, tin or lead, has been modified by the use of so-called "inhibitors" or "restrainers" sold under trade names such as Galvene, Rhodine, Picklette, etc. The effect of the addition of a small percentage of an "inhibitor" to hydrochloric acid is to depress the rate of solution of metal without affecting the rate of solution of the film of oxide (scale). See Evaluation of Pickling Inhibitors, Pirak and Wenzel, Chem.-Ztg. 1932, 56, 193, 214.

Liquid hydrogen ehloride may be stored in steel containers if the gas has been thoroughly dried. The low specific gravity of the liquid and the great weight of the cylinder militate against commercial development of this product.

CHLORISOL. Sodium hypochlorite solu-

tion.

CHLOROCRUORIN. A pigment related to hemoglobin which occurs in the blood plasma of certain polycheto worms (H. M. Fox, Proc. Roy. Soc. 1926, [B], 99, 199).

TRICHLORO-CHLOROFORM, METHANE, CHCl3. Chloroform was discovered by Liebig (Pogg. Ann. 1831, 23, 242) and independently about the same time by Soubeiran (Ann. Chim. Phys. 1831, [ii], 48, 131). Its constitution was determined by Dumas (Ann. Chim. Phys. 1834, [ii], 56, 115), to whom the name chloreform is also due. Chloroform is very widely employed as a general anæsthetic and is an important solvent finding application in the manufacture of fats, rubber, resins, alkaloids and other products. It is a useful preservative.

Chloroform may be obtained by elilorination of a number of organic substances such as alcohol, acetone, acetaldehyde, acetophenone, turpentine, terpenes, etc., but only a few of these are of practical utility.

Chloreform was originally manufactured from ethyl alcohol, but for many years past this has been replaced by acctone. Owing to the much cheapened production of neetaldehyde at the present time chlorination of the latter product bids fair to displace the use of acctone. According to Feyer (Z. Elektrochem. 1919, 25, 115) the formation of chloroform from alcohol by the action of bleaching powder is represented by the

the manufacture of ammonium chloride, both in three stages involving oxidation and forma-

(1) CH, CH, OH+HOCI =CH.·CHO+H.O+HCl

(2)
$$CH_3 \cdot CHO + 3HOC1$$

= $CHCI_3 + H \cdot COOH + 2H_2O$

(3) H·COOH+HOCI=CO2+H2O+HCl

The action of bleaching powder on acetone appears to be best represented by the equation

$$2CO(CH_3)_2 + 6CaOCl_2$$

= $2CHCl_3 + (CH_3 \cdot COO)_2Ca + 2Ca(OH)_2 + 3CaCl_2$

Manufacture of Chloroform from Acetone and Bleaching-powder.—This is the process most generally employed. The method differs in minor details with the various manufacturers, but the following may be taken as representative. The reaction is carried out in a cast-iron still of about 800 gallons capacity, which provided with stirring gear, steam-eoils, and eooling coils, and is connected with a condenser; 300 gallons of water are run into the still, and 800 lb. of bleaching powder are added through a manhole, which is then securely bolted down. During addition of the bleaching powder the mixture is very thoroughly stirred. (In some processes the mixing is carried out in a separate vessel, and the suspension is strained from tho larger unbroken lumps of bleaching powder before being allowed to run into the still.) The container (A in the diagram shown on p. 78) is charged with 70 lb. of acctone, which is then slowly run into the bottom of the still by means of the valvo B. The introduction of the acetone is accompanied by a riso of temperature which is not allowed to exceed 110°F., cooling being effected if necessary by stopping the flow of acctone and circulating cold water through the eooling coil in the still. When all the acetone has been introduced the contents of the still are raised to 134°F. At this temperature chloro-form begins to distil over. The temperature is then very gradually raised to 150°F., so as to keep the chloroform steadily distilling. Towards the end of the reaction the mixture is stirred and the temperature raised until no more chloroform distils over.

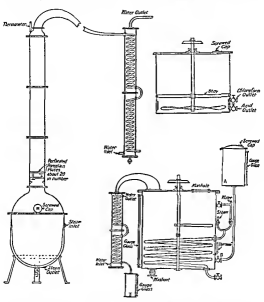
The erude chloroform obtained is separated and purified first by agitation with concentrated sulphurie acid. This operation is carried out in the vessel shown in the diagram; 1,500 lb. of crude ehloroform are introduced into the vessel and thoroughly stirred, by means of the agitating gear shown, with 600 lb. of sulphuric acid. The stirring is continued until a sample of the chloroform when thoroughly shaken with pure concentrated sulphuric acid does not impart the slightest colour to the latter. The time required for complete purification is usually about 3 hours. The chloroform is next separated from the sulphuric acid and finally distilled over lime. The yield obtained from the above quantities averaged from over 2,000 batches was 124 lb., the highest yield in any one case being 131 lb. Variation in yield is attributed to the varying composition of the bleaching powder, though following equations, the reaction taking place doubtless other factors influence the result.

Bleaching powder centaining less than 33% solution of actalochyde with a solution of available chloring rise unsatisfactory results, hypochlorite. A yield of 80% of chloroform a while samples containing more than 35% of clasmed for the following process (G.P. 347450), chlorine are also often unsatisfactory. The best A 20% agreem solution of a cetaldchyde is results appear to be obtained with bleaching allowed to flow into a solution of Sheaching chlorine.

sists essentially of the treatment of an aqueous addition is complete stirring is continued for a

powder containing about 34% of available powder containing 93 g. of active chlorine lorine.

per litre. The mixture is vigorously stirred and Manufacture from Acetaldehyde.—This con. the temperature maintained at 45°C. After



short time, the solution neutralised, and the vided with a surrer and maintained at a tempera-chloroform separated by distillation. Purificature of 70°-80°C. The chloroform is removed G.P. 339914 and F.P. 521700). A continuous into a separating ressel, the chloroform distilling process for the production of chloroform, in off through a condenser while the residual injury which a matture of acctone and acctalablyde is are continuously discharged. treated with a solution of hypochlorite contain.

Other Methods of Manufacture.—Chloroform ing 40 g. of active chlorine or more per later, has been manufactured in America from carbon described in U.S.P. 1915334. The mixture of tetrachloride by reduction with pascent hydro-

cation is effected in the usual way (see also by leading steam into the mixture as it passes

the solutions is led into a reaction vessel pro- gen. The most usual method is to stir vigorously

iron in water, the reaction being initiated by the addition of a small amount of hydrochloric acid (U.S.P. 753325, 1107025). Numerous attempts have also been made to manufacture chloroform hy chlorination of methane (natural gas), but so far without any pronounced success. A number of patents have also dealt with electrolytic methods of production from alcohol or acetone, but very little appears to have been made in this way.

Chloroform is a heavy colourless liquid with a characteristic odour. It has h.p. 61-2° (Baskerville and Hamor, Ind. Eng. Chem. 1912, 4, 278; Wade and Finnemore, J.C.S. 1904, 85, 946), f.p.-64-19° (Keyes et al. J.C.S. 1923, 124, II, 376), d4 1.4989 (Baskerville and Hamor, l.c.).

It is not inflammable hut hurns with a greenish smoky flame when mixed with alcohol and the mixture ignited. It is slightly soluble in water (0.5%) and miscible with most organic solvents.
When a primary aminc is heated with

chloroform in an alcoholic solution of potassium bydroxide an isoeyanide is formed:

$$CHCl_3+RNH_2+3KOH$$

=RNC+3KCl+3H₂O

The reaction employing aniline as the primary amine affords a ready method of detecting chloroform in considerable dilution (1-5,000 to 6,000) the characteristic odour of phenyl iso-eyanide being easily recognised. The reaction is, however, not specific, heing given hy hromo-form, iodoform, chloral, etc. Chloroform also reduces Fehling's solution.

Annesthetic Chloroform.—Chloroform for annesthetie purposes is required to he of a very high standard of purity and the various pharma-coposias detail the tests for a number of impurities which may occur in the product and render it unsuitable for use as an anæsthetic. Puro elileroform under the influence of light, air, and moisture is decomposed more or less rapidly according to conditions, the main products of decomposition being carbonyl ehloride (phosgene) and hydrochloric acid, according to tho

equations:

$CHCl_3+H_2O+O_2=COCl_2+HCl+H_2O_2$ $CHCl_3+H_2O_2=COCl_2+HCl+H_3O$

The presence of a small amount of ethyl alcohol in the chloroform has been shown to exert a very definite stabilising effect, and most pharmacopeins specify the addition of alcohol for this purpose. The amount permitted by the "British Pharmacopeia" is from 1-2%. According to Baskervillo and Hamor (Ind. Eng. Chem. 1912, 4, 368, etc.) alcohol appears to act by reason of its property as a reducing agent, oxidation of alcohol taking place in preference to that of chloroform. At the same time the products of oxidation appear to exert a retarding action and eliloroform containing the amount of alcohol mentioned above, when properly packed and stored so that the action of light and air is reduced to a minimum, will keep in a satisfactory condition for years. The theory of the action of alcohol is supported by the fact that

a mixture of the tetrachloride and finely divided | inhihit the decomposition of chloroform. The action of such substances for the preservation of chloroform for anaesthetic purposes has been investigated by Bodendorf (Apoth.-Ztg. 1929, 44, 351) by exposing chloroform containing the suggested preservative to the action of light from a quartz lamp, when it was conclusively shown that alcohol is the most satisfactory stabilising agent.

For the estimation of chloroform in small amounts W. H. Cole (J. Biol. Chem. 1926, 71, 173) has suggested a colorimetric method based on the colour reaction obtained with pyridine and caustic soda. The method is said to be eapable of detecting 0.001% of chloroform in solution, but its usofor the determination of the eompound in biological fluids has been criticised by Yeager (Proc. Soc. Exp. Biol. Med. 1929, 26, 403). It has, however, heen modified for the purpose by Gettler and Blume (Arch. Path. 1931, 11, 554). A colorimetric method for tho estimation of ehloroform in alcoholie solutious, tinetures, etc., has been described by Moffitt (Analyst, 1933, 58, 2). It depends upon the production of a hlue colour at the ordinary temperature with β -naphthol and potassium hydroxide, a reaction which readily lends itself to colorimetric comparison. A similar colour is given with a naphthol, but in this case the reaction is also given by other chlorinated compounds. With β -naphthol, however, closely related compounds such as earhon tetrachloride, dichloroothylcne, ethylene chloride, etc., givo no colour. A. J. E.

CHLOROGENIC ACID (I), Caffetannio Acid (II). The substance described as (I) hy Payen (Ann. Chim. 1849, [iii], 26, 108) is now recognised to be the chief constituent of Rochleder's (II) (Annalcn, 1846, 59, 300, and many later authors). The alcoholic extract of Liherian eoffee yielded on evaporation 3.3% of crystalline potassium-caffeine chlorogenate, which loses caffeine when shaken with moist chloroform; dilute sulphurio acid then liberates (I), $C_{16}H_{18}O_{9}$, needles, with $\frac{1}{2}H_{2}O$, m.p. 208°, [a]_D -35·2° (Gorter, Annalen, 1908, 358, 327; 1911, 379, 111; Freudenberg, Ber. 1920, 53, [B], 232). Tannase from Aspergillus niger splits the acid into quinie acid and caffeie acid, indicating a depside formula (Freudenberg, l.c.). The position of the linkago was determined by H. O. L. Fischer and Gerda Dangsehat (Ber. 1932, 65, [B], 1037), who assigned to (I) the constitution 3-[3:4-dihydroxycinnamoyl]-1-quinic acid,

Gorter isolated (I) from eoffec leaves, the latex of Castilloa elastica (Rec. trav. chim. 1912, 31, 281), sunflower seeds (Arch. Pharm. 1909, 247, 436), and from Strychnos nux romica (ibid. idem. 197). If Gorter's colour renetion (ibid. 247, 187; Annalen, 1911, 379, 111) could be trusted, (I) is widely distributed in the vegetable kingdom many other organic reducing substances also (cf. Charaux, J. Pharm. Chim. 1910, [vii], 2,

292 ; van der Haar, Pharm, Weelblad, 1920, | methyl alcohol to remove xanthophyll. Finally 57, 194) Caffeie acid gives the same colors-tion (Freudenberg, 1c.) The estimation in [1] in acctone and methyl alcohol with water; this coffee has been studied (B, 1934, 40, 119, 379, 426, 524, 700, 779, 985). Inspite of the name [11], (1) does not precipitate a dilute gelatin solution, but when warmed with a 10% solution gives a precipitate on cooling (Freudenberg, Ic.) J. N. C

CHLOROPHYLL. The colouring matter of green leaves. A mixture of two closely allied substances—chlorophyll a, C₅₅H₂₂O₅N₄Mg, and chlorophyll b, C₅₅H₂₀O₅N₄Mg.

It occurs widely distributed in plants, mainly in the leaves and green stems, in the chloroplasts, together with the yellow (carotenoid) pigments carotene and xanthophy il It is present in both green and brown algo, the latter of which also contain another carotenoid, fucoxanthin It is the active catalyst in the photosynthesis of augars from earbon dioxide and water, and is commercial product. A water-soluble pigment therefore of fundamental importance. It is the only enzyme of known chemical structure

For the bulk of our knowledge of chlorophyll. we are indebted to the hrilliant investigations of

collaborators

Historical — (See Willstätter and Stoll, "Un-tersuchungen uber Chlorophyll," Berlin, 1913 ¹, Marchlewski, "Chemio der Chlorophylle," Braunschweig, 1909) Of early attempts to isolate the green leaf pigment, those of Berzehus (Annalen, 1838, 27, 296) and Verdell (Compt rend, 1851, 33, 699) may be mentioned. The latter believed that it contained mon, like tho harmoglobin of blood, an idea which persisted Other erroncous conceptions for some time which complicated early work were that the pigment contained phosphorus and potassium Valuable exploratory work was subsequently carried out by Schunck (Proc Rey, Soc. 1885, 38, 336, etc.), and Schunck and Marchlewska (Annales 1894, 278, 329) It was thought at one time that samples of chlorophyll isolated from different sources and at different seasons were not identical. This was disproved by Willstatter, whose investigations (1906-1913) mann (Helv. Chim. Acta, 1933, 16, 739) led to a general clarification of the subject. More examined with great thoroughness by Hans Fischer (1929-1936) with the result that the structure is now certain in main outline and only e few details remain to be settled Chlorophyll has not yet been synthesised, although progress remains to be settled. Extraction -Both fresh and dried green leaves

can be used as sources of ehlorophyll The common nettle, first used by Stoles (1852), is a by acid fractionation (see below). convenient source, being cheap, rich in chlorophyll, readily dried while maintaining its colour obteined by acctone extraction is a dark green well, and being poor in enzymes. Extraction wavy solid. Pure chlorophyll a is obtained is best carried out at room temperature with microerystalling by adding petroleum to its acetone containing 15-20% of water (Will-ethereal solution, Slow evaporation of a solution statter). The acetone solution is treated with in a mixture of these solvents gives thin lanceolight petroleum and extracted with water to late leaflets. Chlorophyll b can be obtained remove the bulk of the acctone and then with similarly in a microcrystalline form. In powder

precipitates the chlorophyll, which is filtered off by means of a column of tale. The tale is mashed with petroleum (to remove all carotene and the chloropbyll extracted with ether. The ethereal solution is dried by means of sodium sulphate and the chlorophyll precipitated by the addition of petroleum. (For details see Will-statter, "Chlorophyll," Chapters 3 and 4) The extraction with alcohol, which is used for certain technical purposes, leads to a chemical change (see below).

A commercial chlorophyll is manufactured for the colouring of soap, oils and perfumes. For this purpose green leaves (nettles) are extracted with etbyl alcohol, the solvent distilled and the residue extracted with benzene. The residue from this second extraction is the basis of the This material has a yellowish green shade; to ohtam a hlush green, 1% of copper sulphate is added in the original alcoholic extraction. The Richard Willstatter and Hans Fischer, and their product presumably contains copper in place of the magnesium atom present in the chlorophyll moleculo (see belou)

The chlorophyll content of dried nettle leaves is about 0.8% and about 80% of this (6.5 g, from I kg of leaves) can ho extracted by Willstatter's

procedure.

The chlorophyll so obtained is a mixture of the a and b components, which are present in land plants in the proportion of about 3 . 1. In brown alga the proportion of the b form is much smaller. In spite of claims to the contrary, it is probable that only two forms of chlorophyl exist (Winterstein and Schon, Z. physiol Chem. 1934, 230, 139, but see Conant and Dietz, Nature, 1933, 131, 131; J. Amer Chem, Soc. 1933, 55, 839). Chlorophyll-b can be removed from a petroleum solution of the mixed pigment by extraction with 90% methy! alcohol (Willstatter and Isler, Annalen, 1912, 390, 269), s method which has been improved by Stoll and Wiede Another method of separation is by chromato recently, the chemistry of chlorophyll has been graphic adsorption on a column of calcium carbonate or preferably, powdered augar (Tswett, Ber. 1908, 41, 1352; Winterstein and Stem, Z physiol Chem. 1933, 220, 263) lly this method the two forms of chlorophyll can be separated from one another and from the carotenin this direction has been made. The exact role ords which accompany them in the plant. In of the pigment in the photosynthetic process the examination of structure it is usual to separate not the two forms of chlorophyll itself, but the corresponding degradation products, the pheophorhides, which are easily separated

Physical Properties -The mixed chlorophyll t Referred to henceforward se Willstätter, form chlorophyll a appears blurah black, it gives "Chlorophyll."

powder of the b form is very dark green, the two components of chlorophyll have the Chlorophyll a is coluble in other, absolute ethyl formulæ: alcohol, acetone, chloroform, carbon disulphide and benzene; moderately soluble in cold methyl alcohol, more soluble in hot; very sparingly soluble in light petroleum, but the solubility is raised considerably by the addition of a little alcohol. It is practically insoluble in alcohols containing 20% of water. The solutions are bhish-green with a red fluorescence. The solubility of chlorophyll-b resembles that of the -a component, but is slightly lower throughout The difference is most marked in the ease of light petroleum. The solutions are yellowgreen to green with a brownish-red fluorescence.

Chlorophyll-a sinters and melts between 117° and 120°; chlorophyll b sinters between 86° and 92° and melts to a thick liquid between

120° and 130°.

Absorption Spectra .- Both components of chlorophyll have complex absorption spectra. The main areas of transmission are in the region of 500mµ (green) and 700mµ (red). The main bands in the spectrum of chlorophyll-a have maxima at $662m\mu$ and $433m\mu$; there are also strong bands at $612m\mu$ and $458m\mu$ and weaker bands between the green and orange-red. The end-absorption commences at about 415mu. The absorption spectrum of chlorophyll-b differs from this principally in having the main red band split into two and in the deepening of red band spit into two and in the deepening of the band in the blue at 457m μ , which becomes the strongest band. For further details, see Willstatter, "Chlorophyll," p. 169; Hans Fischer, Oppenheimer's "Handbueli der Bio-ehemie," 1923, p. 351; Conant and Kamerling, J. Amer. Chan. Soc. 1931, 53, 3522; and a series of papers by Stern, Wenderlein and their collaborators. Z. physical Chem. (A), 1034 eollaborators, Z. physikal. Chem. (A), 1934, 170, 337; 1935, 174, 81, 321; 1936, 175, 405; 176, 81; 177, 40, 165, 365, 387.

Optical Activity.—It was thought for a long

time that chlorophy ll was optically mactive, but in 1933 Stoll and Wiedemann (Helv. Chim. Acta, 1933, 16, 307) using red light $(\lambda, 720m\mu)$ found that both chlorophyll a and -b were levorotatory with $[a]_{72}^{15}$, about -265° . The phytyl- and magnesium-free derivatives, the phorbides (see below), were also active. Fischer and Stern have confirmed and extended these observations (Annalca, 1935, 519, 58, 88) There is a difference between these workers' observations on the case of racemisation of the solutions. The porphyrms derived from chlorophyll (see below) are all optically mactive.

Reactions and Structure. - As stated above, pheophorbide (IV):

ehlorophyll
$$a$$
, $C_{55}H_{72}O_5N_4Mg$ ehlorophyll- b , $C_{55}H_{70}O_6N_4Mg$

Chlorophyll-b contains one more atom of oxygen and two less of hydrogen than chlorophyll-a; actually it contains an aldeliyde group in place

of a methyl group.

"Crystalline chlorophyll," chlorophyllides .-It was found by Willstatter and Utzinger (Annalen, 1911, 382, 129) that when ethyl alcohol was used in place of acetone to extract green leaves, the pigment was isolated as bluish-green erystals. This involves a chemical change catalysed by an enzyme, chlorophyllase, present in the leaf. The change is shown below for the -a eomponent

$$C_{34}H_{33}O_3N_4Mg\cdot CO_2C_{20}H_{39}+C_2H_5OH$$
 $Chlorophy ll-a$
 $\rightarrow C_{34}H_{33}O_3N_4Mg\cdot CO_2C_2H_5+C_{20}H_{39}OH$

The compound (I), ethyl chlorophyllide, is tho main constituent of crystalline chlorophyll. The other reaction product is phytyl alcohol, a wax-like, unsaturated primary alcohol. The structure of this has been proved to bo

both by degradation and synthesis (F. G. Fiselier and Lowenberg, Annalen, 1929, 475,

A similar enzymatic alcoholysis with methyl aleohol yields methyl chlorophyllide,

It was found by Willstätter, Hoelieder and Hug (Annalen, 1909, 371, 1) that the same phytyl alcohol was obtained by the decomposition of the ehlorophyll from a large number of different plants. It is the presence of the phytyl group in the molecule which enables chlorophyll to form colloidal solutions.

Phaophorbides (Willstätter and Hocheder, Annalen, 1907, 354, 205).—Hydrated oxalic acid in alcohol leads to a straightforward elimination of magnesium from ehlorophyll. The product is an olive-brown solid, phæopbytin (II). Similarly, treatment of methyl chlorophyllide with a 17% solution of hydrochloric acid in ether leads to the formation of methyl phreophorbide (III) If stronger acids are used, the plu tyl or methyl esters are also hydrolysed with the formation of the corresponding free acid,

The above reactions are shown for the a series, derivatives of chlosophyll, particularly chloric, In actual practice a minimum of physophorbulevas into a series of perophysis. These are, and 3 are obtained. These can be separated by crystalline solids with low acid numbers, and partition between ether and hydrochlore acid are based on the common structural unit, as follows :

Acid Separation (Willstätter, "Chlorophyll," p. 262).—This is a useful general method for the separation of substances of similar structure by taking advantage of small differences in basicity. The mixture is dissolved in ether and extracted with hydrochloric acid in gradually increasing strength. The acid number of the anbstance is the percentage concentration of hydrochloric acid which extracts two thirds of the substance from an equal volume of its ethereal solution. For example, methyl phwophorbides-a and -b have acid numbers of 16 and 21 respectively. The former is completely extracted by 18% acid, the latter requires 23% acid. Extraction of an ethereal solution of a mixture with 17% acid leads to an almost complete separation. The procedure eannot be used with compounds containing magnesium, such as chlorophyll stackf, or the chlorophyllides, unless the chmination of metal is immaterial

From this point the structure and reactions of the more fully studied -a component of chlorophyll and its degradation products will be dealt 1th exclusively.

Chlorin-e -Two of the five oxygen atoms of chlorophyll a aro in a carboxyl group which carries the phytyl group. Two other oxygen atoms are present in the form of a carbomethoxyl group. This is comparatively hard to hydrolyse by acids, but if phaophorbide-s is boiled very rapidly with methyl alcoholio potash for 30 seconds a tribasio acid chlorin e is obtained. This yields a characteristic trimethyl ester (Willstatter and Utzinger, Annalen, 1911, 382, 171). The change involves a hydrolysis of the stable carbomethoxyl group and a conversion of the fifth oxygen into a carboxyl group;

The implication is that the fifth oxygen atom is part of a pseudo-acidic group. From H. Fischer's work this is a cyclic 8 ketonic ester grouping It was previously thought to be a lactone group Essentially the same change has been brought about by the conversion of methyl phæophorbide-a to the trimethyl ester of chlorin e by th action of diazomethane in methyl alroho (11. Fischer and co-workers, Annales, 1932, 499 194 ; 1933, 506, 107).

lly drolysts of ehlorophyll-a stself with potas under non oxidising conditions gives a similar reaction, but in this case the product is th potassium salt of isochlorophyllin-a,

This is converted into chlorin-e hy treatmen with acids, the atom of magnesium being ex changed for two of hydrogen.

Drastic Degradation by Alkalis-Alkalis a high temperatures convert the magnesium-free

porphin (V):

The conventional lettering and numbering of the porphin group is shown above, and a convenient abbreviation in Va. Porphin and the porphyrins (which are its derivatives, substituted in the numbered positions) possess a characteristic type of absorption spectrum, and the power to form stable derivatives with metals, in which the two imino hydrogen atoms are substituted, eg, hy a divalent metal atom. The porphin nucleus exhibits a resemblance to the aromatic type (see Lanstead, Chem. Soc. Annual Rep. 1935, 32, 359).

The porphyrins isolated from chlorophyll derivatives are tabulated below. They all contain the following substituents; hydrogen atoms on the α, β, and δ carbon stoms, methyl groups on the 1, 3, 5, and 8 carbon stoms, ethyl groups on the 2 and 4 carbon atoms. The other substituent groups are indicated in the table :

PORPHYRING PROM CHLOROPHYLL DERIVATIVES

Substituents

9	Name and formula		CH' CH'	CO ₂ H).
al S		6-	7-	۳.
٦				
h	Rhodoporphyrin, C ₃₀ H ₃₁ N ₄ (CO ₂ H),	со,н	p	Н
ī	Phylloporphynn.	100,11	P	
1	CHN.(CO.H)	н	p	Me
-1	Pytroporphyrin, C ₁₉ H ₁₈ N ₄ (CO ₁ H)	н	р	н
t	Phyllogetroporphyrin, CarH, N	н	Et	Me
ŧ	Pyrrostioporphyrin, C ₃₀ H ₃₁ N ₄	н	Et	н

Pyrrox tioporphyrin, the last-named and simplest of these compounds, was at one time believed to be identical with the oxygen-free ætioporphyrin isolated by degradation of the hæmin of blood (ætioporphyrin-III). Although this is not the case, the substances are very closely related, the ætioporphyrin from the blood pigment having an ethyl group in place of a hydrogen atom on C₆. All these structures have been confirmed by synthesis (H. Fischer and coworkers, Annalen, 1928, 461, 221; 1929, 473, 211; 475, 241; 1930, 480, 109, 189; 482, 232). Pyrroporphyrin from chlorophyll has been converted into mesoporphyrin (VI), a degradation product of hæmin.

Drastic Degradation to Pyrroles.—Derivatives of chlorophyll can be converted into simple pyrrole derivatives by two methods, involving oxidation and reduction respectively. Marchlewski (J. pr. Chem. 1902, [ii], 65, 161) oxidised phylloporphyrin with chromic acid to the anhydrido (VII). Willstatter and Asahina (Annalen, 1910, 373, 227) oxidised rhodo-, phyllo-, and pyrro-porphyrins and chlorin-e to a mixture of methylethyl malcinimide (VIII) and hematic acid (IX, i.e. the imide corresponding to VII).

For recent work on this subject, see Fischer and

Breitner (Annalen, 1936, 522, 151).
Willstätter and Asahina (Annalen, 1911, 385, 188) reduced phylloporphyrin with hydrogen iodide and acetic acid to a mixture of alkyl pyrroles containing hæmopyrrole (X), kryptopyrrole (XI), and phyllopyrrole (XII):

The facts enumerated above lead to the following skeleton structure (XIII) for chlorophyll-a:

XIII. (Doublo bonds and pyrrole carbon atoms omitted for clarity.)

The magnesium atom is attached to nitrogen atoms because it is known that oxygen-free compounds, such as the attoporphyrins, can form magnesium derivatives. The carbon atom on C_{γ} is the source of the γ -nicthyl group of phylloporphyrin, and the CO group on C_6 is the source of the 6-earboxyl group in rhodoporphyrin.

The full formula for chlorophyll-a, proposed by H. Fischer (Annalen, 1935, 520, 88), is as follows:

XIV. (Phwophorbide-a has the same formula with H, H for Mg, and H for phytyl.)

It is not possible in the present article to give the complete evidence justifying the various structural features of this formula, other than those already discussed. Some of the more important points are discussed below, and references are given to the original memoirs. The reader is also referred to the general bibliography at the end of the article.

(1) The general arrangement is the same as that of the skeleton structure (XIII). Fischer's

formula (XIV) is not, however, based on a been established by Fischer by analytical simple porphin structure, because the original methods (Annalea, 1931, 485, 1; 486, 107) and double bond between C₂ and C₂ is saturated ¹ by synthesis (bid. 490, 91; 1932, 494, 80; Chlorophy II is therefore a dihy droporphin derivative, as first suggested by Conant (Conant and Kamerling, I.c., Conant, Chow and Dietz, J Amer. Chem Soc. 1934. 56, 2185) These saturated asymmetric carbon atoms are the seat of the optical activity. In a revent paper (Annalen, 1936, 525, 259) Fischer and Lautsch have succeeded in removing the two " extra" hydrogen atoms quantitatively by means of silver oxide or silver acetate in acetic acid

(2) Carbon atoms 6 and y are connected to form a cyclo pentane ring. This ring contains a pseudo acidic & ketonic ester grouping, in which is the carbomethoxyl group which, as has already been mentioned, resists hydrolysis. It is this ring which is opened when chlorin e is formed from pheophorbide a The presence of the keto group was proved independently by Fischer et al (Annalen, 1934, 503, 224) and Stoll and Wiedemann (Helv Chim Acta, 1934, 17, 163) by the preparation of an exime from methyl phrophorbide a The presence of the cyclopentane ring is deduced from Fischer's work on phaoporphyrin a,

(Annalen, 1929, 474, 65, 1930, 478, 54, 284, 480, 197) This substance is prepared by reducing phrophorbide-a to a leuco compound by means of hydrodic and acetic acids, followed by aerial oxidation. The whole process is an somene change, involving the reduction of the vinyl group on C_s, followed by the removal of the two "extra " hydrogen stoms on C_s and C_s. Physoporphyrin a; easily loses its carbo-methoryl group by ketonic fission to yield phyllocrythrin (XV, R=H).

This substance (Marchlewski, Z physiol Chem 1929, 185, 8) is also formed by the biological degradation of chlorophyll, and can be isolated from the frees of runmants. Its structure has

³ The periods of the road Hilberth Indicates atom. By period at C₀ and C₁ above, cannot as yet be exceeded at an extention for a presided affect represents to discovered the control of the periods of the control of the con

497, 181).

(3) The reasons for placing the phytyl group in the propionic acid side-chain, (C₇), and hence the methyl group on the carboxyl adjacent to C, are given by Conant and Hyde (J. Amer. Chem Soc 1929, 51, 3668; 1931, 53, 2382, cf. Fischer, Sus, and Klebs, Annalen, 1931, 490, 38).

(4) The substituent group on C, in chlorophyll and the phorbides is vinyl (and not ethyl. as in the corresponding purphy rins). It reacts additively (i) with diazoacetic ester (Fischer and Medick, Annalen, 1935, 517, 245) and (ii) with cold by driodic acid in a stream of oxygen (" oxoreaction," Fischer and Riedmair, Annalen, 1933, 505, 87) The vinyl groups of the phorbides undergo the following changes in these reactions:

The Phase Test (Molisch)-This is a colour reaction given by both forms of chlorophyll, and by derivatives such as the phorbides, phyllides, and chloring, in which the dihydroporphin (phorbin) arrangement is maintained. It is not given by porphyrins An ethereal solution of chlorophyll or a derivative is shaken with methyl alcoholic potash The colour changes from green to bright yellow-brown and then back to green The reaction is essentially an aerial oxidation, consuming two equivalents of oxygen (Steele, J Amer Chem Soc. 1931, 53, 3171) A similar reaction occurs when an alcoholic solution of chlorophyll is evaporated to dryness or allowed to stand in contact with air. The chlorophyll is said to be allowerised and loses its power of ervatallisation (Willstätter and Utzinger, Annaien, 1911, 382, 129). This was first recognised as an oxidation by Conant et al. (J. Amer Chem, Soc 1931, 53, 359, 2382). The products formed in these reactions have been studied by Conant and by Fascher, but their chemistry is complex

(cf. Chem Soc. Annual Rep. 1935, 32, 385).

Partial Synthesis of Chlorophyll—As has already been mentioned, Fischer and his coworkers have aynthesised phyllogrythm, (XV, R=H) For a complete synthesis of chloro phyll it is necessar; (i) to convert this into phacoporphy in a_1 (XV, $R = CO_2Me$); (ii) to isomerise this to phecophorbide a_1 (iii) to introduce the phytylgroup into the side chain on C7; and (s.) to introduce the central magnesium atom into the phrophy tin so formed The last two stages have been achieved by Fischer et al (Annales, 1934, 510, 156; 1935, 519, 244); the remainder of the synthetic problem awaits solution.

Chlorophyll b -This differs from chlorophyll-d

Bibliography—R Willstatter and A. Stoll, "Untersuchingen über Chlorophyll," Berlin, 1913; Il Lischer in Oppenheimer's "Handbuch de Biocheme, 1923, p. 351, 1930, 87, 1933, 262, J.C.S. 1934, 245, R.P. Linstend, Chem. Soc. Annual Rep. 1935, 32, 362; 1937, 34, 375; for the occurrence of substances of the chlorophyll series in organic minerals, see A. Treibs, Angew Chem 1936, 49, 682

RPL

CHLOROPHYLLASE 19 an enzyme of the esterase class discovered by Willstatter and Stoll (Annalen, 1910, 378, 18, 1911, 380, 148), which in ethyl alcoholic solution splits off phytol from chlorophyll, forming cthyl chlorophyllide The enzyme is widely distributed in leaves remains in the leaf residues after the chlorophyll has been extracted with alcohol. The method of preparation has been improved by Norck (Biochem Z 1927, 183, 135) It is inactivated in the leaf powder by dialysis, but restored by the addition of CaCla

ЕГА

CHLOROPRENE 1 BUTADIENES AND Poly of Frins (conjugate d)

CHLOROXYL. Cinchophen by drochloride Quinophan hydrochloride Cinchophen is 2 plicingly quinoline 1 carboxylic acid,

C.H. C.H.N COOH

(Eli Lilly, London), B P C.

CHLOROXYLONINE, C22H23O7N, an alkaloid occurring in East Indian satishwood (I'r bors satishe) (Chloroxylon swietenia DC Rutacere) It crystallises from EtOH in colourless prisms, in p 182°-183°, [a]_D¹⁸ -9° 18' (in chloroform), is soluble in excess of ammonia The salts are solution and is neutral to litmus erystalline and dissociate in H₂O, anrichloride, B HCl AuCl₃, groups of reddish yellow needles, in p. 70°. Chloroxylonine contains four methoxyl but no hydroxyl groups (Anid, JCS 1909, 95, 961) It produces derinatitis when applied to the skin and is the cause of this trouble sometimes experienced by satinwood workers (Cash, Brit Med J. 1911, Oct 7, and Bull Inip Inst 1911, 9, 351).

CHLORPICRIN or CHLOROPICRIN. (Trichloronitromethane), CCl₃ NO₂, is formed by the action of aqua regia on many organic compounds, eg cymene, styrene, etc (Ontta and Fernandes J. Amer Chem Soc. 1916, 38, 1813), on acctone or the by-products of the acctone industry (Boyd, J S C I 1925, 44T, 222), and by the action of nitrie acid alone on many aliphatic chloro compounds such as chloral and chloroform (Dănăili and Soare, Bul Soc Romann Stunt, 1932, 35, 53). It is most readily prepared by the action of chlorinating and oxidising agents on organic nitro compounds, thus Green and Rowe obtained chloropierin by treating nitro p phenylenedianime, dinitranilme, intro access p phenylenediamine, etc., with sodium hypochlorite. The oxidation of picric acid or its salts with chlorine and bisic materials num hypochlorite. The oxidation of pierie acid 14, 251. Complete summary 1848-1932, United or its salts with chlorine and basic materials States, Dept of Agric., Miscellaneous Publito yield chloropierin forms the subject of many cations, No. 176.

methyl on C₃ (formula XIV), but the structure patents differing chiefly in the nature of the has not been conclusively established basic constituent which may be a metallic oxide, carbonate, or borate (B P 142878), hme (U S P. 1413198; see also Frahm, Rec trav clim 1931, 50, 1125), or bleaching powder (U S P. 1327714, 1996388), etc

Chloropierin is a mobile oil, mp -69°C, bp. 112 C, of great chemical activity, solutions in organic solvents often depositing aminonium chloride in the light (Pintti, Gazzetta, 1921, 51, 1, 145) or yielding other products by photo-chemical reaction (Alexejewski, Chem Zentr. Reducing agents readily attack 1933, 1, 3683) eliloropierin, iron and hydrochloric acid giving a theoretical yield of methylamine (Frankland et al , JCS 1919, 115, 159) while merenptans are reduced with formation of disulphides (Nekrassow and Melnikow, Ber 1929, 62, [B], 2091). Grignard reagents effect replacement of the three chlorine atoms by alkyl or aryl radicals necompanied by luminescence (Wedland Z. Physik, 1906, 4, 417, Asing and Dufford, Physical Rev. 1933, [2], 44, 315)

Chioropierin is best recognised by the colour reaction with dimethylaniline paper or by the formation of mitrite on reduction with metallic ealcum (Alevejewski, J. Chem. Ind. Russ. 1931, 8, 50). Quantitative determination in the atmosphere is effected either by absorption in acctic neid and reduction with iron, or by thermal decomposition over potassium earbonite at 350°, the chlorine in both methods being estimated gravimetrically (Deckert, Z. Hyg

1929, 109, 485)

Chloropierin is highly poisonous (toleration limit for man, 60 cu mm per cu m. air (Flury, Z ges exp Med 1921, 13, 567), eausing lachrymation, coughing, vomiting and eventually death by overwhelming edema of the lings It has a corrosive action on the skin and frequently produces abscesses (Underhill, Arch Int Med 1919, 23, 753), exposure of wounds, ents, ete, to its action invariably producing ulceration. The lethal properties of chloropicrin have led to its wide use as a disinfectant (F P. 677340), preservativo (Plucker, Z Unters Lebensm. 1932, 63, 313), insecticide, fungicide, fumigant, etc. Its ready detection recommends it as a warning agent in illuminating gas (G P. 447655) and in hydrogen cyanide (Zentr. Gewerbeling u Unfallverhutung, 1927, 14, [3], 65, USP. 1949466) It has found many other uses, among which may be mentioned its use in vetermary practice (Cliem. Rep. US Dept Agr. 1924, 12), the purification of vaccines (1.P. 612075), and the extermination of rats, foxes and other pests (Compt rend Acad Agric France, 1921, 7, 568) Among purely chemical uses may be mentioned its employment in the manufacture of crystal violet (U.S.P. 1402195) and as an oxidising agent in the synthesis of quinoline and derivatives (B.P. 198462) Chloro-pierin has been used in gas warfare and is regarded as the typical gas of the non persistent deadly type (1 CHEMICAL WARFARF)

Literature.- Review of important work up to and including 1931, Jackson, Chem. Rev. 1934,

CHLORQUINOL, 2-chloro-1.4-dihydroxy- | porated in plain or milk chocolate, to give nut

B.P.C

CHOCOLATE, from the Mexican word " Chocolatl," is the term applied to certain types of cocoa preparations intended for human conaumption, either in the form of a beverage or as a confection. No legal atandards have been adopted in the United Kingdom, although proposals to this effect have been made (e.g. R. Whymper, Congress of Cocoa and Chorolate Makers, Berne, 1911, and N. P. Booth, Seventh International Congress of Applied Chemistry, 1909). The majority of chocolate preparations contain augar, but some, usually prepared for special purposes, are unsweetened or have saccharin as the sweetening agent. Those to be consumed in the form of beverage often contain a rertain proportion, about 20%, of arrowroot or some other farmaceous substance. They may be in block form or as finely ground powder. The larger proportion of chocolate, however, is in the form of confectionery and thus also falls into well-defined classes

Manufacture — The most simple type is what is usually known as plain chocolate, consisting eventually of cocoa and sugar. The cocoa, in the form of block cocoa from fermented beans (see Cucoa), is first prepared by granding in a mill between granite or steel rollers at a tempera ture (approximately 35°-40°C) sufficiently high to melt the cocoa butter present and to reduce the whole to liquid form. In order to give a product which is amouth to the taste, the particles of fat free cocoa must be ground as finely as possible. Mason (Analyst, 1933, 58, 440) states that in well known proprietary brands the maximum length of the particles varied from I to 200 The sugar used in the manufacture of good class chocolate consists of white high grade crystals, while in lower quality chocolates an inferior grade is sometimes need. It ahould be dried and ground to a fine powder and may, with advantage, be warmed pre viously to admixture, in order to maintain the temperature of the mass In special cases where sugar is madmissible, as ni chocolate intended for use by diabetic subjects, the sugar is replaced by saccharm It is then usual, in order to give the necessary body to the chocolate, to add some form of meal or flour (preferably leguminous) and pea, bean, or linseed meal are preferred for this purpose Flavourings, chiefly vanilla, but less frequently einnamon, closes, mice, nutureg or their essential oils, are almost invariably added, usually in alcoholic solution In the cheap forms of plum chocolate substances are used which many regard as adulterants. Chief among these are wheat or potato flour, rice, arrowroot, dextrin and ground cocoa shell Milk chocolate is prepared by including in the molten mass a certain quantity of dried milk powder Although of course, the proportions used by different makers vary greatly, the following may be regarded as a typical composition: 15% fat free cocoa matter, 25% cocoa butter, 46% ugar, and 20% milk powder

benzene, mp. 106°, hp. 263°. chocolate and nut milk chocolate. All these CHLORYLEN. A brand of trichlorethylene. forms of confectionery chocolate are run, whilst Used as an antineuralgic (Schering, London). still in a molten condition, into moulds, giving, on cooling, bars, blocks, cakes and numerous fancy chapes.

Many types of chocolate confectionery are manufactured and sold, in which the chocolate forms an envelope or covering for other substances such as creams, biscuits, dried fruits, nnts, fruit jelly, hqueurs, and medicinal preparations such as pills The most important of these fancy preparations are the creams. The interior cream is made either of powdered sugar or a mixture of this with glucose, with or without flavouring and colouring materials. The mixture is heated until it melts and, while in the liquid state, is poured into starch moulds of the desired chape on a table which can be agitated hy mechanical means. On cooling, the creams set into a stiff granular mass. They are then coated with the hot liquid chocolate paste. In order to render the paste more mobile when heated and to give smoothness to the finished article, additional fat is added to the chocolate, This may be either cocca butter or a substitute fat such as illipé. The covering is effected either by dipping the creams in the paste or by pouring the latter over the creams, uniformity being increased by means of the rocking table In recent years machinery has been devised, and so now in use, to carry out the covering peocess.

Chemical Examination -The chemical ex ammation of chocolate follows well defined lines, the principal features bring the extraction, determination, and examination of the fat, the determination of the percentage of sugar, and a microscopical examination of the fat-free residue which will indicate the necessity for the determination, if present, of cocoa shell or furinaccous matter. In the case of chocolate covered goods the covering and interior are separated, the proportions determined, and the two parts then

examined independently. Owing to the extensive use of substitutes for cocoa butter the complete extraction of the fat, not only for the determination of the proportion, hut also for its subsequent examination, is a matter of considerable importance. The Southet method, using light petroleum as a solvent, takes considerable time, and various methods have been suggested, for all of which rapidity, without loss of accuracy, is claimed. Bloomberg suggests extraction for 4 hours with anhydrous ether In support of this, Fincke (Bull Off-intern fahr, choc. 1932, 2, 327) states that, if there is a possibility of a substitute fit being present, ether is preferable to light petroleum, as the latter will not dissolve out the way which is likely to be present in the adulterant fat. Lepper and Waterman (J. Assoc Off. Ague-Chem. 1925, 8, 705), however, state that preliminary drying is necessary and, further, that the method is inaccurate owing to the extraction of theobromine and other substances They suggest a method of exhausting the sample by several extractions with light petroleum, collecting the extract in a tared Erlenmeyer flash, and cocco butter, 40% augar, and 20% milk powder ung the extract in a tared Erlenmeyer flash, and Nuts, chiefly hazel or almond, are often moor exaporating the solvent, drying, and weighing

1919, 119, 104) suggests that the sample should be subjected to two preliminary washings with 50% alcohol, the mixture being centrifuged and the liquid decanted off. The process is then repeated several times with a mixture of equal volumes of ether and light petroleum, the suceessive ethereal layers being decanted off into a tared flask, the solvents evaporated and the fat dried and weighed. In order to avoid the evaporation of the solvent and the weighing of the dried residue, Hasso and Bake (Chem.-Ztg. 1923, 47, 557) suggest the extraction of the fat by a mixture of equal volumes of ether and methyl ethyl ether. The difference between the respective refractometer readings of the solution and of the solvent is referred to tables which give the appropriate percentage of fat. Ruffy (Mitt. Lebensni, Hyg. 1926, 17, 75) has attempted, with some slight modification, to apply the Gerber method of estimating fat in milk, cheese and meat. The presence of sugar in the chocolato presents a complication on account of its carbonisation by the sulphuric acid. It is claimed that this has been overcome by treating 3.5 g. of the chocolate with 20 ml. of other in a strong walled tube. The mixture is well slinken and centrifuged; 10 ml. of the clear liquid can then be transferred by a pipette to a hutyrometer containing 13 ml. of sulphurie acid (density 1.5). This is shaken vigorously, centrifuged for 5 minutes, and the reading taken. Hartel and Jacger (Z. Nahr. Genussm. 1922, 44, 291) suggest that the total fat should be extracted from milk chocolate with light petroleum and its Reichert Meissl value determined. value, multiplied by the percentage of fat and divided by thirty, gives the proportion of milk fat, which is taken to be one-quarter of the whole milk solids. In a later communication, however, Hürtel (ibid. 1924, 48, 32) states that if a certain kind of milk powder has been used, it is impossible to obtain all the fat by extraction with ether or light petroleum. In considering the fat extracted from milk chocolate, the possibility that skimmed milk has been used should not be ignored. The butyrie and saponification numbers afford a reliable and rapid indication of this.

The proportion of sugar present can readily be determined by the polarimeter, after clarifying the solution with basic lead acetate. For accuracy in this method von Fellenberg and Ruffy (Mitt. Lebensm. Hyg. 1932, 23, 6) state that certain precantions, which they describe, are essential, owing to the disturbing effect of the fat-free cocoa matter. Macara and Hinton (Congrès. Intern. Fabr. Chocolat. et Cacao, 1930, 341) have made an extensive investigation of the subject and evaluated the various known methods for plain and for milk chocolate. The examination of the latter presents special difficulty, owing to the varying and uncertain character of the milk used in manufacture. Hartel and Jaeger (l.c.) give a method in which the amount of milk solids is determined from the content of milk fat (v. supra). Whilst the determination of the lactose must be made on the original substance, the proportion of easein ean

in the usual manner. Hughes (Chem. News, | with sodium earbonate, as described in the paper. This is elaborated by Beythien and Pannwitz (Z. Nahr.-Genussm. 1923, 46, 223) who state that the ratio of easein to lactose in twenty milk chocolates varied from 1.16 to 2.74. A method of determination of the milk content of chocolate, based upon the proportion of CaO in the ash, is suggested by Grossfeld (ibid. 1922, 44,

The presence of lecithin in cocoa and its preparations is of technical, and therefore of chemical, importance. Although the reason is not clear, the presence of lecithin promotes tho mobility of hot eocoa or chocolate mass, but this is not so marked when milk has been added. Nottbohm and Mayer (Z. Unters. Lebensm. 1933, 65, 55) have stated that free leeithin is absent from cacao beans but that cholino from phosphatides, other than leeithin, may occur in the alcoholic extract, possibly produced by the action of the hot alcohol or by the reasting process. It is generally accepted, however, that some lecithin is present in the cocoa mass in varying proportions under 1%. The proportion is sometimes increased by the addition of commercial lecithin, thus making possible the extraction of an additional 5% of fat without loss of mobility in the hot mass. Various methods for the determination of leeithin have been proposed, one being the Soxhlet extraction with alcohol, the solvent being evaporated off and the residue treated with 30% H₂O₂ and supplierie acid. The solution is then neutralised with ammonia and the phosphate from the leeithin precipitated with a mixture of strychnine nitrate and ammonium molybdate. The precipitate is filtered, dried and weighed. Winkler and Sale (J. Assoc. Off. Agrie, Chem. 1931, 14, 537) after extracting with alcohol and light petroleum, tako up the dried extract with obloroform, add alcoholie potash, and evaporate again to dryness. The residue is then taken up in nitric acid and the P₂O₅ determined. Avent and Morgan (J.S.C.I. 1932, 51, 169) consider that the best solvent to use in the extraction is a mixture consisting of four parts of benzene and one of alcohol.

If the microscopic examination of the fat-free residue indicates the presence of foreign starch or shell, the former can be estimated by the ordinary diastase method, allowance being made in the calculation of the result for approximately 14% of starch in the fat-free cocoa material. although it is better to make an independent determination of this value. The proportion of shell can be estimated by determination of the crude fibre, allowing for 6.10% in the fatfree rib and 18.25% in the fat-free shell, although in this case also an independent determination of the standards is to be advised. Offitt (d. Assoc. Off. Agric. Chem. 1932, 15, 546) suggests certain modifications which render the fibre method applicable to milk chocolate. ("The Chemistry, Flavouring and Manufacture of Chorolate Confectionery and Cocoa ") quite properly emphasises the point that the fibre determination alone is probably only positive when over 25% of shell is present. A more decisive analytical interpretation may be placed be determined by treating the fat-free dry residue lupon the ratio of the nitrogen to fibre which

differs very considerably for nib and husk; | butters" or "green butters" are used, and, fibre nitrogen for the former being 0 98-1-27.

The dietetic value of chocolate has often been emphasised and has been the subject of conaderable discussion Knapp ("Cocca and Chocolate," 1920), quoting from the Journal of the Royal Agricultural Society of 1914, states that the "food value" of the cacao bean is represented by 183 units as compared with turnips 8, carrots 12, potatoes 26, nee 102, wheat 106, peas 113, and oatmeal 117 He also quotes the United States Department of Agriculture to the effect that the fuel value in Ih, calories of cocos (0 5 oz to 1 pint of water) is 65 compared with tea (0.5 oz per pint) 15, and coffce (1.0 oz per pint) 16. The figures represent the products without milk or sugar He states that I pint of cocoa made of one third milk, and of cocos, and I or of sugar would have a fuel value of 320 calories, and is therefore equivalent, in this respect, to 1 lb of beef or 4 eggs Knapp further quotea data showing that eating chocolate has 2,538 calonea per lb compared with sugar 1,815, catmeal 1,811, bread 1,180 beef ateak 960, eggs 594, milk 314, potatoes 302, apples 214, and cod fish 209 Sasaki and Wakayama (J. Agric Chem Soc Japan, 1932, 8, 160) found that plain chocolate contained antineuritie vitamin B, 3 g of choco late per day being enough to cure antineuritie vitamin deficiency in pigeons Jensen (le). however, considers that the vitamin content of ordinary plain chorolate is unimportant, but the consumption of good class milk chocolate ensures valuable calcium and phosphorus assimi lation by the definite vitamin D activity Vitamin A is also present in proportion which increases the food value of milk chocolate

F. G H. T CHOCOLATE FATS (Confectionery Fats) I nder this name may be grouped those fats which are employed in the manufacture of chocolate and confectionery, for which purposes a non greasy, digestible fat, which displays e peculiar hard and brittle consistency (" snap ") together with a relatively low melting-point (not by cacao-butter (q e), which is both the typical and most prized fat of this class, and is employed for the highest grades of chocolate, etc., but for cheaper products, cacao butter may be replaced, either wholly or in part. by cheaper substitutes Animal fats, which were used in earlier days, are liable to impart an unpleasant " ammal flavour " to the choco late, and at present only edible fats of vegetable origin are employed. Among the most impor organ are employed. Among rus most smpor tags. The presence do some work of tast of these are Borneo (disple) fallow (g. seems to be seems tal to the life of all cells, but as (which, apart from its funtly greensh colour, jet little is known as to the part they ply clockly results) and command an excellent aubstituted exception of certain invertebrates, cholesterol is and the refined and decolored "steams" present and appears to be synthesised in time. prepared from coconut and palm-kernel oils propries from coconit and pain-zeroet one placefroe and arigmaneroe occur of the hyperparts of the hyp sometimes shea fat and other "regetable In 1775, cholesterol was discovered by Conradi

more recently, bydrogenated coconut and palmaverage 12, and for the latter 53-60, average kernel oils; the latter are harder than the corresponding natural fats, and are also claimed to assist in the prevention of "sugar bloom" of chocolates. The use of hydrogenated oils, such as arachis, cottonseed or sunflower-seed oils, of appropriate melting point has been proposed, but as a rule the texture of such products is not satisfactory The use of Dika fat (q v., from the seed kernels of species of Irringia), which possesses a suitable melting point and "snap" has also been suggested from time to time (cf. Baudon, Bull. Mat Grasses, 1926, 10, 38).

It may be noted that the definitions for

chocolate products adopted (the English delegation dissenting) by the International Congress of Manufacturers of Coron and Chocolate at Antwerp (1930) intentionally imply a prohibition of the use in "chocolate" of fats other than expressed cacao butter the veto also extends to the use of solvent extracted cacao fat or the fat from the germs and shell of the cacao bean. Similar standards have been adopted or are under consideration in the food legislation of many countries, including Austrie, Belgium, Germany, Italy, the United States, Switzerland, France, New Zealand and South Africa, which in general forbids the undeclared inclusion in chocolate products of all fats other than expressed cacao butter (apart, of course, from the meidental introduction of milk fat or nut-fats associated with the corresponding ingredients of milk or nut chocolates) The Imperial Economic Com-mittee in its Report on "Cocoa" (22nd Rept. 88/503,22, 1932, HMSO) has recommended that analogous legislation forbidding the use of more than a small proportion of cacao butter aubstitutes in the manufacture of chorolate pastes, etc. should be enacted in the British Empiro, the provision of a small margin for other fats as suggested in accordance with the amendment of the English delegates to the Antworp Congress, in order to allow the nonfraudulent inclusion of a holesome materials which might genuinely improve the product Up to the present (1938) no legislative steps have been taken to implement the Committee's over 37°-38°C) is required This particular recommendations in the United Kingdom, where combination of properties is well exemplified the chocolate manufacturers regard auch regulations as undestrable.

E L elloCHOLANIC ACIO r. CARDIAC GLYCO-SIDES (Vol II, p 383a).

CHOLELYSE. A preparation of oleic acid for cholchthama.

CHOLESTEROL is a member of a group of naturally occurring unsaturated secondary alcohols, containing a nucleus of four reduced rings. The presence of some one of these sterols present and appears to be synthesised in tiro. Sitosterol and sigmasterol occur in plant relia

stituent. It occurs either free or as its fatty acid esters in all animal tissues; brain and nerve tissue, suprarenal glands and egg yolk are particularly rich in it. It is readily extracted from gallstones; it may be prepared by grinding up sheep's brain or other animal raw material with sand and plaster of Paris, or the mineed tissue may be dried at 100°, ground in a coffee mill and the mass extracted with ether or acetone. The residue from the extract, after removing the solvent, is saponified with alcoholic potash, the alcohol distilled off and the eholesterol separated by extraction with ether. The formation of emultions may be largely avoided by saponifying the other extract of the tissue with an alcoholic solution of sodium ethoxide at ordinary tem-After filtering off the compact perature. inass of soaps which separates, the cholesterol erystallises out on concentrating the alcoholether filtrate.

Properties.—Cholesterol is insoluble in water, that advanced by Rosenbein sparingly soluble in cold, easily in boiling alcohol and Ind. 1932, 51, 454, 954).

in gall-stones, of which it forms the chief con- | (sp.gr. 0.87), and in CHCl3, ether, CS2, less soluble in acetone, petroleum, benzene and glacial acetie acid. It is soluble in fats, fatty acids, and in solutions of bile salts. It erystallises from ether or CHCl₃ in silky needles, in.p. 150° (corr.), from 90% alcohol it forms characteristic plates, containing one molecule water of crystallisation. Sp.gr. 1.046; optically active; [a]p -31.12° in ether solution, -36.61° in CHCl3. Sublimes unchanged at 200°, but is decomposed at higher temperatures, hydrogen being evolved at 300°-320°. Distils unchanged in vacuo.

Constitution .- Our knowledge of the structure of cholesterol is based on a long series of researches initiated by Mauthner in 1893 and carried out by, amongst others, Diels, Borsche, and especially by Windaus and his co-workers from 1903-1932. The formula which now finds general acceptance is a modification by Wieland and Dane (Z. physiol. Chem. 1932, 210, 268) of that advanced by Rosenheim and King (Chem.

The connection with the bile acids first suggested by Lifschutz (Ber. 1914, 47, 1459) was definitely established by Windaus and Neukirchen (Ber. 1919, 52, [B], 1915); the researches of Wicland and his colleagues on the structure of the bile acids have led to the identification of a series of degradation products, common both to cholesterol and the bile acids, which has been of the greatest importance in establishing their constitutions. The similarity of the structure of these substances with that of the group of sexual hormones is also of physiological importance.

Hydrogenation.—Reduction with platinum and hydrogen gives cholestanol (dilivdrocholesterol), m.p. 142° (Willstätter and Mayer, Ber. 1908, 41, 2199), of which coprostanol (coprosterol) occurring in the intestines is an isomeride (Dorec and Gardner, J.C.S. 1908, 93, 1625). Reduction with nickel and hydrogen at 200° gave a mixture of dihydroderivatives, δ. and ε-cholestanols; δ-cholestanol on boiling with amyl alcohol and sodium amyl oxide gave eoprosterol, separated as the digitonide and recovered from this by boiling in xylene (Windaus, Ber. 1916, 49, 1724). δ- and ε-Cholestanols were shown to be the isomerie forms of cholestanol and coprostanol respectively which depend on the inversion of the OH group. Neither δ- nor c-cholestanol gives a precipitate with digitonin and they should now be known as epicholestanol and epicoprostanol respectively.

Oxidation.-Various oxidising agents, c.g. nitrie and chromic acids, have produced a long series of oxidation products some of which are referred to below. By the action of ozone an ozonide containing 7 atoms O is formed (Dorée and Gardner, J.C.S. 1908, 93, 1328; 1909, 95, 638; 1916, 109, 46).

The Presence of a Secondary Alcohol Group was proved since the ketone cholestenone (m.p. 78°) was formed by oxidation with CuO at 300° (Diels and Abderhalden, Ber. 1904, 37, 3092) or by oxidation of cholesteryl bromide with chromic acid in acetic solution at 70° (Windaus, Ber. 1906, 36, 518) and removal of the Br by zine and acetic acid; in the latter reaction Tschesche has shown that the ethenoid link in cholestenone has moved from the 5:6 to the 4:5 position.

The Position of the Ethenoid and Secondary Alcohol Groups.—Originally Windaus postulated the presence of a terminal vinyl group; if, however, cholesterol is oxidised with fuming nitrie acid (Windaus, Ber. 1903, 36, 3752; 1917, 50, 133) and the unsaturated nitro-derivative obtained treated with zine and acctic acid, nitrogen is removed as ammonia and the keto-alcohol, cholestanonol, $C_{22}H_{16}O_{2}$ (m.p. 140°), is formed, which on further oxidation with chromic acid gives cholestanedione (m.p. 170°-171°); this may also be obtained by oxidising cholesterol with cold chromic acid or KOBr solution to cholestantriol, oxidising further to cholestanonediol, splitting off water and reducing the

and reduction yields a C₁₇ dibasic acid which on __CH OH groups must also be separated by at further oxidation is converted to a tetrabasic acid least two earbon atoms (Windaus and von containing 27 C atoms, and which does not lose Staden, Ber. 1921, 54, [B], 1059).

cholestenedione thus formed (Windaus and CO, on heating. The carbinol and ethepoid Kirchner, Ber. 1920, 53, [B], 614). If cholestanonol is converted to the corre-sponding chloroketone, the latter on oxidation the asme carbon atom, the -CH CH- and

Structure of the Side-chain -Oxidation of Idibasie acid produced was converted to an aubycholesterol with nitric acid led to the isolation dride and by applying Blano's rule that a 6 carbon dinitrosopropane, and acetic, succinie, methyl euccinic, and a glutaric acids, acetone, hydroxywahutyne acid, methyl wohexylketone and wooctane, each of which may be regarded as derived from an 8-carbon side chain having the atructure -CHMe [CH₂]₂ CHMe₂. Since acetone and cholamo acid are formed by the oxidation of coprostane, which is known to contain the same side-chain as cholesterol, this structure for the side chain was confirmed when Wieland, Jacobi. and Schlichting (Z. physiol. Chem 1926, 161, 80), by means of alternate Grignard condensations and oxidations, removed successive carbon atoms from the aide-chain of cholame acid until finally etiocholanone.

(mp. 105°), was left, the ketone group being present in ring IV, to which the side chain was attached.

Ring Structure of the Nucleus -Up to 1931 the presence of two reduced rings had been established in the cholesterol molecule hydrocholesterol on oxidation gave the dibasie acid C27H44O4, which on heating lost CO, and water, giving a cyclic ketone, C22H44O, and this on further oxidation gave a C22 dibasic send, which lost water on heating, forming a cyclic Ring II, was converted through its nitro-denvative to a ketone and the latter oxidused, the cholesteryl chloride, a mixture of hydrocarbons

of a series of products amongst which Windaus chain dibasio acid (e.g. adipie) on heating gives a (Z. physiol. Chem. 1918, 162, 160; Windaus cyclic ketone and a 5-carbon dibase acid (e.g. and Resau, Ber. 1913, 46, 1245) identified glutaric), an analydrade, a5 carbon ring structure. glutaric), an anhydride, a 5 carbon ring atructure was attributed to Bing II. The same reasoning led to a 5-carbon ring structure being later attri-bated to Ring IV. The applicability of the Bland rule to the determination of ring atructure was criticised by Rosenheim and King (Chem-and Ind 1932, 51, 464) and on other grounds a 6 carbon ring structure is now attributed to both Rings I and II. Evidence that Rings III and IV are present in the nucleus of cholestered was supplied by the work of Wieland and his colleagues on the bile acids (q.r.). By bacterial reduction of cholesterol in the intestines, coprosterol, isomene with dihydrocholesterol, formed. This gives on reduction pseudocholestane (isomeric with cholestane) which on oxidetion yields a cholsnic acid identical with that addangs from the chole and or dule. The three hydroxyl groups of the latter may be reduced giving C₂₄H₄₀O₂, cholanic acid. The saturated hydrocarbon, cholestane, C₂₇H₄₂, obtained by complete reduction of cholesterol (Mauthor, Monatsh. 1907, 23, 1113; 1909, 30, 635) yields on oxidation actions and crystalines cholane acid, C₂,H₄₀O₃ (Windaws and Newkirchen, Ber. 1919, 52, [B], 1915) isomerie with bile cholane acid. The empirical formula C₂₇H₄₈ for the saturated hydrocarbons cholenges. stane and coprostane, itself indicated the presence of a 4 ring system (Diels and Linn, Ber. 1908, 41, 544). Forther knowledge as to the structure of the aleleton of the sterol nucleus was supplied anhydride. To ring 1, the refore, by Blane's rule a by the work of Diels et al. (Ber. 1927, 60, [B]. 140. cyclohexane strocture was assigned (Windaus Annaien, 1927, 459, 1910 nh dedrydogenation and Dalmer, Ber 1919, 52, [B], 162). When Products of cholestene, containing the ensaturated link in Mauther and Sudd (Monatheft, 1806, 17, 29) obtained as thermal decomposition products of

tained the 8-carbon side-chain of cholesterol; from the dehydrogenation products of this compound by hydrogen and selenium at 240°-310°, Diels isolated chrysenc. Cholcsterol similarly treated gave chryseno and two hydrocarbons, C₁₈H₁₆ and C₂₅H₂₁. The presence of four rings in the nucleus was therefore established. Tho hydrocarbon C₁₈H₁₆ was identified by Rosenheim and King and by Kon as 3-methylcyclopentenophenauthrene. In chrysene, the Me group at position 13 in cholesterol has entered the 5-carbon ring IV forming a 6-carbon ring. C₂,H₂₄ was identified as a pentacyclic hydro-carbon, formed by ring closure between the terminal side-chain and the nucleus, thus giving evidence as to the position of attachment of the side-chain. The grouping

must, therefore, he present in the cholesterol molecole.

When it had been established that the nucleus contained 17 C atoms, the side-chain 8 C atoms, 2 C atoms remain to be accounted for. Evidence has been given above that a methyl group occurs at position 13, and the other methyl group is placed at position 10 for the following reasons, when ring I of the cholesterol molecule was broken down step by step to a monobasic acid by the action of Grignard's reagent this acid was very difficult to esterify indicating that a tertiary carbon atom containing a methyl group was probably adjacent to the COOH group. For this and other reasons position 10 has been assigned to the second methyl group.

X-ray examination of the sterols by Bernal (Chem, and Ind. 1932, 51, 466) showed that the previously accepted formulæ did not accord with the crystallographic measurements, whereas Rosenheim and King's formula (ibid. 464) is supported both by the crystallographic measure-

ments and the optical evidence.

Isomerism in the Cholesterol Series .- A large number of isomeric derivatives exist. Structural isomerides arise from variation in the position of the double bond. Stercoisomerides due to the inversion of the OH group occur and roay be differentiated by the action of digitonin, those containing the OH group in the same position as cholesterol being precipitated by digitonin, no such precipitate being formed when the OH group is in the inverted position. Stereoisomerism due to the relative position of Rings I and II also takes place. Thus cholesterol and cholestene may be converted to coprosterol and coprostene by the action of HCl. Cholestane and its derivatives are regarded as the trans-derivatives; the coprostanc series as the cis-compounds. Windans (Ber. 1904, 37, 4753) pre-pared isomeric oxyketodibasic acids, C₂₇H₄₂O₆, determined by Hübl's method if the conditions one forming a lactonic acid (m.p. 192°-193') are closely defined; Wijs reagent gives variable

of the formula C19H28, which no longer con- which on heating with potash gave an oxyketo acid incapable of lactonisation (trans form), and concluded that the reduced ring structure must be capable of cis-trans isomerism.

Nomenclature.—Considerable confusion exists in the nomenclature of the cholesterol derivatives since names were given before the relationships of the various compounds bad been elucidated. It has been suggested by Rosenheim and King (Chem. and Ird. 1934, 53, 91) that certain useful simplifications showing structural connections should now be made. The term pseudo has been used to denote both sterie and structural isomerism; it seems desirable now to use the term "epi" for all substances formed by steric inversion of the OH group, e.g. epicholestanol, cpicoprostanol, these substances do not give insoluble digitonides. The term copro should be given to the cis- derivatives related to coprostcrol; some of the changes involved would be:

Old nomenelature. (C15- derivatives.)	New nomenclature. (Cis- derivatives.)
Coprosterol (saturated alcohol) Allocholesterol (unsaturated alcobol) Coprostanone Coprostenone Pseudocholestane Pseudocholestene	Coprostanol Coprostanone Coprostanone Coprostanone Coprostane Cholestene

Cholesteryl Esters .- Those of the lower fatty acids are prepared by heating cholesterol with the anhydrides of the acids; those of the higher fatty acids by boiling the cholesterol with the acid and a dehydrating agent. These substances on melting form anisotropic liquids, many of the esters having three stable liquid The acetate melts at 114°, the propionale at 98°. The benzoale melts at 144°-145° to a turbid liquid which examined with tho polariscope shows definito crystal structure and at 178° passes to a clear liquid, the p-nitrobenzoate melts at 189°, p-aminobenzoate at 238°. The laurate can be heated several degrees above its melting-point without melting (Shriner and Ko, J. Biol. Chem. 1928, 80, 1).

Detection .- Cholesterol crystallises in characteristic thin rhombic plates from 90% alcohol. Under the microscope they often show notched angles and exhibit a curious overlapping arrangement. A number of colour tests are given by the unsaturated sterols; the most important for cholesterol is Salkowski's test, in which 2 e.e. of a CHCl2 solution of cholesterol are shaken with an equal volume of eone. H2SO4, the CHCl3 layer passes through blood-red and cherry-red to purple, which lasts for several days; poured into a basin, it becomes successively blue, green and yellow owing to absorption of water. The sulphuric acid layer is strongly fluorescent.

Iodine Value.—This may be satisfactorily

and much too high results, owing either to laterol compound. The cholesterol is recovered substitution or, according to Schoenheimer, by heating the prespitate with 100 cc glacul because re formation of the double bond table sectie and go. Z. dust for 2 hours, the excess of place with liberation of soline (J. Biol. Chem. zmc filtered off, and a large quantity of water 1935, 110, 461) The re-in ecids also give abnormally high values with Wijs solution. The enthracene hydrocarbons react similarly with Hubl and Wijs reagents, whereas the phenanthrenes give much higher values with Wijs than with the Huhl solution (Smedley MacLean to the ease with which, when mixed with ohie and Thomas, Biochem J 1921, 15, 321) Dam recommends a solution of bromme and the preparation of ointments and salves; it is pyridine sulphate (8 g Br, 10 g cone H₂SO₄, and 8 g pyridine per litre) (Biochem Z 1924,

Estimation - The total amount of cholesterol in a ti-sue may be determined by precipitating the alcoholic solution of the uneaponifiable fraction with a 10 solution of digitonin in 9000 alcohol, after standing for some hours, the repeatedly kneading with uater and centric precipitate, containing 25% holesterol, is figure 1 from purified product is atterwards collected on a Gooch filter, washed with alcohol kneaded with 36% water and forms the hydroxy and other, dued at 100°, and weighed. If it is desired to estimate separately the free and about 100% of unter, does not become raned, combined cholesterol, the alcoholic solution of the ether soluble extract must first be pre cipiteted with the digitonin solution and the resulue then asponified end the cholesterol estimated in the unsaponifiable fraction Preceher (Z Nahr Genussm 1917, 33, 491) re commends that after saponification the soap should be decomposed with hydrochloric acid, and the chloroform solution of the fatty acids seperated, filtered and treated with the digitorin solution at 50°, the liquid being stirred during the precipitation. A micro method for the estimation of cholesterol hy oxidation of the digitoride is described by Okey (J Biol Chem 1930, 88, 367) In fata where adulteration with vegeteble oils has taken place, it is important to discriminate between the vegetable and animal sterols, for this the acetates or dibromides must

be prepared and their melting points examined. Marcusson and Schilling recommend the following method (Chem -Ztg 1913, 27, 1001). 50 g fat are shaken for 15 minutes in a separating funnel with 20 ce 1% alcoholic solution of digitonin. The bottom layer of oil is run off, the alcoholic upper layer containing the flocen lent precipitate of digitoride is shaken with 50 to 100 cc ether. The precipitate is washed with ether, dried, powdered, again washed, and then heated with 1 5 e.c. acetic anhydride in a test tube for half an hour On cooling, cholesterol and situaterol acetates ery stallise out. The m p. of the recrystallised acetates are respectively 114° and 128°. Cholesterol and suosterol may also be separated by crystallising their dibromides from other and glacial acctio and (Windaus, Chem Ztg. 1906, 30, 1011) A mixture of 4 g cholesterol and 4 g autosterol 19 dissolved in 80 e c. ether and 80 c c. 5% bromine in glacial acetic added. After standing for one hour at 0°, the precipitate is washed with 40 c.e. glacial acetic acid, and with 40 c.e. 50% acetic aced, the washings being added is formed by the hidrolysis of lecithin and hence to the filtrate and a further precipitate added probably occurs in combination in all living to the original precipitate, which is then washed cells; its isolation from animal and vegetable with water and dried. The precipitate contains tissues has been recorded very many times, but the cholesterol dibromide, the filtrate the sito- has no significance, since it is readily set free

added The sitosterol is obtained by treating camilarly the solution of the situsterol dibromide

Cholesterol occurs with other sterols in a preparation of wool fat known as linobn and used mainly for medicinal purposes; owing oil, it is absorbed by the skin, it is used in also used in the manufacture of cosmetics and pomades and for softening leather. Wool fat, which contains about 70% sterol and 30% fatty acids, is saponified by caustic soda giving an emulsion which on dilution with water gives the so called wool milk. On centrifuging im pure landline separates out and is purified by kneaded with 30% water and forms the hydrons wood fat of the BP Anhydrous lanolin absorb and is antiseptic A commercial preparation of ebolesterol extracted from the apinel cord of

cattle is also evailable 150Cholesterol, m p. 1370-1380, [a]p=+600, also occurs in wool fat and may be separated from cholesterol by heating the mixed sterols for 30 hours in a sealed tube with four times their weight of benzoic acid. The benzoates are in-oluble in boiling alcohol and are separated by ery stallisting from ether Cholestery 1 benzoate erystallises in shining plates, isocholesterol benzoate as a light powder, mp. 190°-191°. asoCholesterol gives no colour reaction with chloroform and sulphuric eeed and is not precipitated by digitonin. When shaken with sulphuric acid, the yellow solution gradually changes to brownish red, in the Liebermann-Burchard test, a brown red colour is obtained Windaus and Tachesche (Z. physiol Chem. 1930, 190, 58) showed that isocholestend is a mixture of two sterols both containing 30 carbon

CHOLESTEROPHAN. Dimethyl para bante acid,

CHOLIC ACID, C24H40O4, 3 7:12 tn

bydroxycholanic acid (r. Bile Acids) AND RELATED SUB-CHOLINE STANCES. Choline, trimethyl · β - hydroxy-ethylammonium hydroxide,

enzyme action. Kauffmann (Z. physiol. Chem. 1911, 74, 175) has shown that when perfectly fresh ox brain is worked up rapidly, with suitable precautions, hydrolysis may be avoided and no choline is obtained. The free choline in semen is the result of enzyme action (Kahane and Levy, Compt. rend. 1936, 202, 2186). The best natural source is crude lecithin; hard egg-yolk is extracted with other and then with alcohol; the extracts are mixed and evaporated and the residue is boiled for 1 hour with 50 parts of cold saturated baryta solution. After removal of the baryta as carbonate or sulphate, the filtrate is evaporated and the residue is extracted absolute alcohol. Alcoholic with mereurie chloride then precipitates choline mercuri-chloride, C₅H₁₄ONCl-6HgCl₂.H₂O, which salt can be crystallised from hot water; it is very slightly soluble in the cold. After removal of the mercury by hydrogen sulphide and evaporation of the filtrate, choline chloride is extracted from the residue by alcohol. For the isolation of small quantities of choline, along with betaine, from plants, see Schulze (Z. physiol, Chem. 1909, 60, 155). The final separation depends on the ready solubility of choline chloride in absolute alcohol, which leaves betaine chloride behind. Choline is, however, best prepared synthetically, by the method of Renshaw (J. Amer. Chem. Soc. 1910, 32, 128); rather more than one equivalent of trimethylamine gas is passed into ethylene elilorohydrin in a tuhe cooled to -12° to -20°, which is subsequently heated to 80°-90°; the yield is almost quantitative. (For another method, starting from ethylene dibromide, see Kriiger and Bergell, Ber. 1903, 36,

Choline can be recognised to some extent qualitatively by means of its diehroitic crystalline periodide (Florence's test for semen; Bocarius, Z. physiol. Chem. 1901, 34, 339; Rosenheim, J. Physiol. 1905–1906, 33, 220). This salt is precipitated at a concentration of choline of 10⁻⁶ by adding a concentrated solution of iodine in potassium iodide. If the choline solution contains sodium biearbonate. the weak base betaine is not precipitated, and can be separated as periodide after acidification of the filtrate from the corresponding choline salt (Stanek, Z. physiol. Chem. 1905, 46, 280; 1906, 47, 83; 48, 334). The reineckate of choline, [Cr(SCN)₄(NH₃)₂]NMe₃C₂H₄·OH, with a solubility of 10⁻⁵ may also be employed qualitatively and quantitatively. The mercurichloride mentioned above forms crossed hexagonal prisms, m.p. 249°-251°, soluble in 50 parts of water at room temperature, but practically insoluble in absolute alcoliol. The platini-chloride, (C₅H₁₄ON)₂PtCl₆, dissolves in 5.8 parts of water at 21° and is dimorphous; the conversion of the regular isotropic into the anisotropic monoclinic form is the surest means of identifying choline without analysis (Kauffmanu and Vorlander, Ber. 1910, 43, 2735). The ourichloride dissolves in 75 parts of water at 21° and in hot alcohol. Neither the gold nor the platinum salt has a characteristic meltingpoint. The free base and many of its salts are

from phosphatides by chemical processes or by enzyme action. Kauffmann (Z. physiol. Chem. 1898, 24, 513). For soluble non-deliquescent salts, see G.P. 290523, 290740. Minute 1911, 74, 175) has shown that when perfectly fresh ox brain is worked up rapidly, with suitable precautions, hydrolysis may be avoided and no choline is obtained. The free choline in semen is the result of enzyme action (Kahane and Levy, ibid. 1916, 77, 408).

HO·NMe3·CH2·CH2·OAc, Acetylcholine, has during the last decade come to be recognised as a substance of great physiological importance. Its powerful action in lowering the blood pressure (many thousand times as great as that of choline) was noticed by Hunt and de Taveau in 1911 (U.S. Hyg. Labor. Bull. 73). It was first recognised in ergot by Dale (J. Physiol. 1914, 48, 3) and isolated by Ewins (Biochem. J. 1914, 8, 44). Dale and Dudley (J. Physiol. 1929, 68, 97) isolated it from the spleen of horses, in which tissue physiological tests indicated the presence of 10 mg. per kg. Bisehoff, Grab and Kapfhammer (Z. physiol. Chem. 1930, 191, 179; 1931, 199, 135; 200, 153) claim to have isolated 28 mg. per litre from ox blood by means of Reinecke salt, but this relatively enormous yield has not been confirmed by others. Acetylcholine is secreted by the endings of the parasympathetic nerves, when the nerves are stimulated, and so acetylcholine is responsible for the humoral transmission of certain nervous impulses; thus it passes into the fluid perfusing a frog's heart, when the vagus nerve is stimulated, and the fluid can then affect a second heart (Loewi). In these experiments the amount of acetylcholine is too small to bo isolated, but its presence may be inferred from its rapid hydrolysis and inactivation by boiling, by alkali, or by an enzyme. This onzyme, choline-esterase, was isolated from horse sernm by Stedman, Stedman and Easson (Biochem. J. 1932, 26, 2056); its action is inhihited by physostigmine and similarly constituted synthetic urethanes. Hence physostigmine poisoning is in reality a poisoning by the subject's own acetylcholine. The synthesis of acetylcholine in ox brain has recently been demonstrated by Stedman and Stedman (Biochem. J. 1937, 31, 817); they isolated 18 mg, as the double eliloroplatinate of choline and acetylcholine from minced ox brains, after 2 hours incubation at 37°; physiological tests (on lecch muscle) indicated that 48 mg, had been formed. The last-named paper and one by Dudley (Biochem. J. 1929, 23, 1064) should be consulted for the difficult technique of isolating this very unstable ester in the presence of much larger quantities of choline.

prisms, u.p. $249^{\circ}-251^{\circ}$, soluble in 50 parts of water at room temperature, but practically insoluble in absolute alcohol. The platinischloride, $(C_5H_{14}ON)_2$ PtCl₆, dissolves in 5-8 parts of water at 21° and is dimorphous; the conversion of the regular isotropic into the anisotropic monoclinic form is the surest means of identifying choline without analysis (Kauffmann and Vorlander, Ber. 1910, 43, 2735). The aurichloride dissolves in 75 parts of water at 21° and in hot alcohol. Neither the gold nor the platinum salt has a characteristic meltingpoint. The free base and many of its salts are syrupy (for details, see Gulewitsch, Z. physiol.

related to that of aretylcholme and, according time. Easson and Stedman (I.c.), who have to Kögl, Dusberg and Eraleben (Annalen, 1931, 489, 156) probably the constitution

HO NMez-CH(CHO)CHEt OH.

The alkaloid encopin in black mustard seeds is the choline ester of smapic acid. In connection with the physiological action of snake venom, Fourneau and Page (i.e.) found polmityl- and s'eard- choline to be strongly hemolytic. Acetyl choline and some related denvatives have been used tentatively in therapeuties; the most im portant of these is perhaps carbaminoylcholine (Doryl of E. Merck) Other sub-tances in which eholine is esterified with phosphoric acid have been isolated from animal tissues (liver); they seem to result from the partial hydrolysis of lecithin and sphyngomyelin; for these and the biochemistry of choline, are Kahane (Bull. Soc. Chim. biol. 1937, 19, 207).

Other toses related to choline are neurone or raultrimethylammonium hydroxide,

CH, CH NMe, OH,

formed from choline by putrefaction and having a similar physiological action, but 10–20 times at tonic. β dynasodisjd deckel or β-laydrary thydanine (edamine), HO CH, CH, MH, was of the briain putreface kephalin (i.e. kenthan without X methyl groups). Colanine has also reviewed the colonic state of the briain putreface kephalin (i.e. kenthan without X methyl groups). Colanine has also reviewed the colonic state of the colon For a fuller account of choline and related sub-For a injer account of choine and relaces sub-etances, see Gurgenbeim, "De biogenen Amine," 2nd ed., Berlin, 1934 (3rd ed. in pre-paration), and for the biochemistry of choline, Kahane, Bull. Soc Chim. brol. 1937, 19, 203.

CHOLINE ESTERASE is an enzyme of the lipace class which hydrolyses acctylcholine, HO NMe, CH, CH OÁC (r. CROLIXE). view of the outstanding importance of acetylcho-Lne as the substance by means of which the parasympathetic nerves produce their effect, existence in the body of an enzyme capable of and mucoids and has been investigated by destroying it is of interest.

It is apparently a specific enzyme and widely distributed in the tissues (Galehr and Plattner, Pflüger's Archiv 1927, 218, 488, 506). A method of preparing it from blood serum is described by Stedman and Stedman (Boochem, J. 1935, 29, 2563). Its activity is best determined by titration with 0-02 N-NaOH with bromothymol blue as indicator (Stedman, Stedman and White, Biochem. J. 1933, 27, 1955). It is destroyed by heat above 56°.

It acts on other estees of cholme; an increase in the size of the scyl group increases the rate of hydrolysis unless a aide chain has been introduced, when a decrease occurs (Easson and Stedman, Proc. Roy. Soc. 1936, B 121, 142).

The order is n-butyry 1>propions 1>isobutyry1

Theorder manonty:

The influence of pg on choine esterage to chrome spinel or picotite; on use some extent depends on the substrate; is some extent depends on the substrate; but optimize the some extent depends on the substrate; but optimize the some extent depends on the substrate; but optimize the some extent depends on the substrate is optimized by ferro conde, there may be a transtount to pure transition the substrate is substrated by ferro conde, there may be a transtount of the summerstal are cube; in expatilisation and before the same isomorphous group—the spinel to extend a mount of chromic ords or the same isomorphous group—the spinel or picotite; on use passage to chrome spinel

most recently studied the question, assume that the substances which inhibit the enzyme act by combining with it in the same way as the choline esters and thus blocking the active groups. In this connection an interesting calculation has been made in regard to the activity of the enzyme, namely, that each active group in it hydrolyses about 1,490 mola. of aretylcholine per second. The amount of inhabitor required to produre a large inhabition of the enzyme is very minute.

E. F. A. CHOLUM or CHOLAM r. CERELLA

MILLEY CHONORIN. An albuminous substance allied to and resembling gelatin, obtained by boiling cartilage with water. It dries to a hard borny substance which softens to a jelly in cold water, and is soluble in boiling water, An aqueous solution is decomposed by 2% aluminsum bydroxide into chondrostan aulphune and (c. Choverents) which remains in solution and has [a]₀ -46 59° and the chondrin residue ([a]₀ -386 85°) which is adsorbed in the collod; this reaction affords a method of differentiating between choudrin and glutin, as a solution of the latter gives an immediate precipitate with barrum chloride, whilst a solution of chorden gives no appreciable precipitate, but if the solution is stirred with 10% aluminium hydroxide and filtered, the filtrato gives with barrum chloride an immediate precipitate of the banum salt of chondroitin sulphume acid (Rakuzin and Braudo, J Russ Phys. Chem. Soc. 1926, 52, 397, Z. Unters Lebensm 1926, 52, 397;

Pakuzm, Chem. Zig 1923, 47, 602) CHONDODENORINE,

ORINE, an ecoquinoline alkaloid. CHONOROITIN 15 a tetrasacchande con sisting of two glururome send and two chondros-In amine units.

E. F. A. CHONOROSAMINE 19 2 aminopalactor. It has been obtained from some of the mucins Levene and La Forge (J. Biol. Chem. 1914, 18, 123 ; 1915, 20, 443).

E. F. A. CHOPIN EXTENSIMETER r. Berad (Vol. II, p. 783). CHOPPING NUT or ESERÉ NUT r

CELABAR BEAN (Vol. II, p. 1984).
CHROMATOGRAPHY t. CAROTENOMS

(Vol II. p. 398) and CHEMICAL ANALYSIS (ibid.,

CHROMITE or CHROME-IRON-ORE, A member of the spinel group of minerals, con austing essentially of ferrous oxide and chromic oxide, FeO Cr.O. or FeCr.O. analogous to the aluminates. The ferrous oxide is, however, often partly replaced by magnesia, and the chromic oxide by alumina, so that there is a analyses; the ordinary ore, as mined, contains on an average 45% Cr2O3.

Cr,O3	Al ₂ O ₃	Fe ₂ O ₃	FeO	MgO
I. 67.9			$32 \cdot 1$	
II. 44·15	$22 \cdot 41$	5.78	11.76	15.67
HI. 59-20	7.15	n.d.	25.02	4.42
IV. 41.23	24.58	2.28	16.99	14.77
V. 56·54	12.13	-	18.01	14.08

I. Calculated for FeCr₂O₄.

II. Franklin, Macon Co., North Carolina.

III. Price's Creek, Yancey Co., North Carolina (J. H. Pratt, Amer. J. Sel. 1809, 7, 281). Also SiO₂ 2-80, MaO 0-69.

IV. Tampadel, Lower Silesia (H. Traube, Z. deut. Geol. Ges. 1894, 46, 50).

V. Dun Mountain, New Zealand.

(For analyses of chromite isolated from meteorites, see W. Tassin, Proc. U.S. Nat. Museum, 1008, 34, 085.)

34, 685.)

In general appearance, chrome-iron-ore is very like magnetite, but it is readily distin-guished from this by the dark-brown colour of its strenk or powder, and hy the fact that it is only slightly, if at all, magnetie. Sp.gr. about 4.5. Crystals are of rare occurrence, the mineral being usually found as grains disseminated in basic rocks of igneous origin, especially those rich in olivine (viz. the peridotites). Sometimes these grains are acgregated into nodular masses of considerable size and with a granular to compact texture. Rocks of this kind, when subjected to weathering processes, become hydrated and altered into serpentine, and for this reason mest of the workable deposits of chrome-ironore are in scrpentine rocks. With the denuda-tien and breaking down of these rocks, the heavy grains of the chemically resisting chromite collect in the beds of streams and rivers; and some of the deposits worked in the Ural Mountains are of this nature.

Chromite is the only commercial source of chromium and its compounds. It is used for the preparation of the oxides and chromates employed as pigments, and in dycing, calico-printing and tanning; in the manufacture of chrome-steel and ferrochrome, and for chromebricks and furnace linings. It is mined in Asia Miner, Serbin, Greece, the Ural Mountains, New Caledonia, India, Baluchistan, Rhodesia, Quebee, Chlifornia and Oregon; and formerly, to a considerable extent, at Baro Hills, near Baltimore in Mnry land, and in Lancaster Co. in Pennsylvania. Quarries were, at one time, werked on Unst, one of the Shetland Isles (Special Reports on the Mineral Resources of Great Britain, Vol. 5, Mem. Geol. Survey, 1916).

References.-Chrome ore and chromium, Imperial Mineral Resources Bureau, London, 1920. On the chemical composition of chromite, see L. W. Fisher, Amer. Min. 1929, 14, 341. On Canadian deposits, F. Cirkel, Dept. of Mines, Canada, No. 29, 1909; and those of Rhodesia, F. E. Keep, Bull. Geol. Survey Southern Rhodesia, 1930, 16.

L. J. S. Chromite is one of the most important neutral refractories, especially in metallurgical furnaces. It is said to have been used in France in 1872

varies considerably, as shown by the following | melting-point, in the neighbourhood of 1,950°C., it is not acted upon hy siliceous fluxes, and for that reason is often placed as a "buffer" layer between the basic magnesia and acid Dinas brick in Siemens-Martin and other furnaces. On account of its resistance to gases, flue-dust and slag particles, chrome bricks are used for the ports of gas-fired furnaces, tap-holes, etc. It is employed for the construction of furnaces for the smelting of copper and nickel ores (Glenn, Eng. and Min. J. 1901, 72, 637; J.S.C.I. 1902, 21, 52), and is also very resistant to oxides of

antimony, lead and bismuth.

Chromite should contain over 40% of chromic oxide and less than 6% of silica, the remainder being composed of ferrous oxide, alumina, lime, and magnesia. In some cases it is employed in the form of lumps or loose powder which can be packed in position and afterwards sintered or even fused into a monolithic layer. The material may also be ground, mixed with a suitable binder, moulded into bricks, which must be fired at a high temperature. It has been stated that clay could not he used as a binder, but chromite mixed with from 10 to 15% of Zettlitz kaolin gave a very refraetory product (Simonis, Stahl u. Eisen, 1908, 28, 334; J.S.C.I. 1908, 27, 283). The most usual binders are refractory clays, hauxite, magnesite and lime, the last giving a highly refractory brick. Chromite bricks are heavy (ep.gr. ahout 4.5), close-textured and fairly strong mechanically, but, like magnesia bricks, fail suddenly at about 1,400°C, under a load of 50 lh. to the sq. in. Wologdine (Bull. Soc. d'Encour. 1909, 108, i, 879; J.S.C.I. 1909, 28, 709) gives the following figures for the heat conductivity:

. 0.0057 C.G.S. units. Chromite I. . 0.0055 agglomerated 0.0028

The electrical resistivity at various temperatures has been measured by Hartmann, Sullivan and Allen (Trans. Electrochem. Soc. 1920, 38, 279) and by McLeod and M'Mahon, working under Professor Stansfield at McGill University (ibid. 1912, 22, 89). The two sets of measurements which differ somewhat are given in the following table:

	I	Iartn	ann et. al.	Star	nsfield e	t. al.
Cold .		48.1	Megohms.			
800°C.	. 8	03 O	hms	2,800 O	hms per	cin. eubo
900°C.	. 5	25 ,		760	••	,,
1,000°C.		71		420	,,	**
1,100°C.		78,	•	430		**
1.200°C.	•	<u>63</u> ,	,	450		,,
1,300°C.	•	77	,	410	,,	**
1,400°C.	•	85 ,	•	320	71	**
1,500°C.		41.	•			

Chromite appears to expand regularly with increasing temperature and has a larger thermal expansion than any other ordinary refractory material, except magnesite, amounting to 1.6% at 1,500°C. (B. Bogitch, Compt. rend. 1921, **173,** 1358).

During the last few years, a considerable amount of attention has been given to the effect of relatively large admixtures of chromite with other refractory materials, especially chronic-silica, chrome-magnesite, and chrome-alumina and earlier in Austria. In addition to a high mixtures. In some cases, possibly most marked

with chrome alumina mixtures, beating to a confirmed, a wheat plant being killed by a high temperature may bring about chemical of 00641% solution of sodium dichromate, whereas reaction between the ovides with the formation; it required a 0.3% solution of chromium sulphate of new compounds, but in others the chromate to give a similar result. The red coloration retains its identity. At the same time, its given by sodium 18 dihydroxy naphthalene presence modifies the hehaviour of the mixture 36-disulphonate was utilised for the identificain service Thus, chrome silica brieks are mechanically stronger at high temperatures than plun chrome bricks, and tend to spall less than either chrome or silica bricks. Similarly, chrome-magnesite bricks have a greater resis tance to spalling than many fireds, bricks and a higher "refractoriness under load" than normal silica bricks (Budnikoff and Mandelgrin, Ber Deut. keram Ges 1932, 13, 527, Lynam and Rees, Trans Ceram, Soc. 1936, 35, 138; sbut 1937, 36, 110 et eq. Chesters and Lee, sbut. 1937, 36, 294). Ricke and Ungewiss, Ber. Deut. Keram Ges 1935, 16, 482)

The following analyses, taken from various sources, illustrate the composition of chromites

for refractories :

	101 79	99 87	98 96	
SIO,	1 87	2 60	5 20	
CaO	5 10	3 07	0.78	
MgO	3 79	1 10	9 22	
Al,O,	3 17	2 60	13 90	
Fe,O,	36 63	28 10	16 20	
Cr _s O ₃	51 23	62 20	53 66	

W C. H CHROMIUM. Symbol Cr, at wt 52 01, at no 24, 1sotopes 50, 52, 53, 54 The common chromium mineral and the only important ore of chromium is chromite, FeCr,O, (q v.) Chromium also occurs in chrome diopside and in the mica known as fuchsite, which is distinguished from other varieties by its brilliant green colour, in chromium garnet and in certain chloritic chromium minerals which accompany the chromite As lead chromate it occurs in crocossile, PhCrO4, and as basic lead chromate in the rare numeral melanchroite or phanicochroite (formerly phanicite) 3PhO 2CrO, Me'eorites often contain chromite but rarely more than Io The red colour of the ruby (Papish and O Leary, Ind Eng Chem [Anal], 1931, 3, 13), the greenish colour of serpentine, verde untico marble, emeralds and many other minerals are due to the presence of oxides of chromium (Sainte-Claire Deville and Carron, Ann Chim 1865, [n], 5, 108, cf. Verneud, Compt. rend 1910, 150, 185). Mitchell (Astrophys J. 1930, 71, 1) confirmed the presence of chromium hues in the solar spectrum Spectrographie analysis has revealed the presence of traces of chromium in the ash of all the organs, somewhat more concentrated in the thyroid and spleen (Dutost and Zbinden, Compt. rend 1930, 190, 172) Demarcay reported the presence of chromium in traces in wood ashes (ibid. 1200, 130, 91).

Koonig (Landw. Jahrh. 1910, 39, 775, JCS 1911, 100, n, 524) described the stimulative and toxic effects of various chromium compounds on plants. The results obtained by other workers, that the higher the degree of oxidation have been prepared in the free state. There are of chromoum the more toxic its effect, were three series of perchromatis, red MaCrO,

tion of traces of chromium in plant ash Work. men exposed to the dust of potassium dichromate have suffered from skin eruptions and inflammation of the septum of the nose There is a risk of chromium poisoning in lithography, founding, tanning, and dyeing where chromium compounds are handled, and also from the spray from chromium plating haths (Dobbs, JSC.I. 1930, 49, 163T). The protection of workers has been studied by the Metropolitan Life Insurance Co. (Service Bur. Bull. No 408; J. Amer. Leather Chem. Assoc. 1936, 31, 110; of Fed. abid, 31, 65, Halle aux Cuirs, 1935, 149) Chromic salts are less toxic than chromates (Akatsuka and Fairhall, J. Ind. 113g 1934, 16, 25; D. Brard, "Toxicologie du chrome," Paris, Hermann & Cic. 1935).

Chromium has many uses in the arts; it forms a series of alloys with iron, carbon, nickel, manganese, etc., which include the stainless steels; it is deposited electrolytically as a bright coating which does not tarmsh, in the process of chromium plating. Chromic salts are used in tanning leather and for the preparation of chromium green. Chromates, usually sodium or potassium dichromate, are also employed in tanning, and to an important extent in wool dyeing and as chromated gelatin in photo-mechanical and colour printing. The siksh chromates are employed in the preparation of the pigments, lead chromate (chrome yellow), and zino chromste (zino yellow) In organic chemistry chromio acid is used as an oxidising agent, chromic oxide serves as a catalist in the synthesis of methyl alcohol and in the cyclisation of hydrocarbons, Cf. Homer Ad kins, "Reactions of Hydrogen with Organic Compounds over Copper Chromium Oxide and Nickel talysts," University of Wisconsin Press, 1937,

The accentific interest of chromium is considerable, the metal can assume a passive state, st forms many series of coloured compounds corresponding to all the valencies from 2 to 6 melusive. Ct11, The blue solutions of chromous salta containing the ion Cri+ are powerful reducing agents. Criii The chromic salts derived from Cr₁O₂ evist in two states, viole crystallisable solutions which on holling become green and do not then crystallise. These solu tions may contain unstable hydrated ions, eg [CrCl 5H,O]++, Cr11. The sellow chromates and orange dichromates derived from the trioxide CrO₃ ionise in solution to CrO₄— and Cr₃O₇—. Cr¹³. Quadrivalent chromium is contained in the series of salts derived from triphen; I chromium hydroxide (C,H,),CrOH Cr. The corresponding tetraphen; I chromium salts (C, H,), CrX contain quinques alent chromium The chromohy drocarbons

(CaHs), Cr and (CaHs), Cr

pounds [(NH₃)₃CrO₄]. A very large number | chromium. of chromammines and analogous substances containing pyridine, urea, etc., have been described.

The ethylene diamine compound [CrCl₂·2en]Cl and also the double exalate K₃[Cr(C₂O₄)₃] thave been separated into optically active forms.

been supplied by the Imperial Institute in The resolution indicates an octahedral structure | Mny, 1938.

blue MCrO6, and brown co-ordination com- | for these and other co-ordination compounds of

For n review including chronite, chromium, and ferroehrome, see Roush, Mineral Ind. 1936,

WORLD'S PRODUCTION OF CHROME ORE. (Long Tons.)

Country.	1930	1931	1932	1933	1934	1935	1936	1937
British Empire Southern Rhodesia . Union of South Africa India Other British Empire Foreign Countries Greece Yugoslavia Cuba Japan Turkey New Caledonia Philippine Islands . Other foreign	202,385 13,508 50,684 1,676 23,032 65,606 50,583 38,380 11,421 27,750 60,916	80,334 22,967 19,913 226 5,545 72,123 57,462 11,882 9,573 24,987 72,979	15,445 19,065 17,865 454 1,531 76,242 38,524 500 12,295 68,332	34,493 33,541 15,526 918 14,550 80,913 25,833 21,837 19,681 74,189 49,281	70,961 60,388 21,576 2,781 30,209 85,943 40,604 57,325 26,792 117,951 54,310	104,240 89,003 39,127 2,795 29,309 181,500 51,540 (b)42,081 35,736 148,096 54,437 (b)1,272	172,896 49,486 1,739 46,599 216,000 53,190	62,307 5,000 55,061 (d) 58,918 (c) 93,098 (d) 189,468 47,264
countries	1,827	3,589	603	3,544	1,133	840	4,364	5,644
Total	548,000	381,000	251,000	374,000	576,000	780,000	1,043,000	(d)

(a) Figures up to and including year 1934 refer to the year ending Sept. 30th; figures for 1935 and 1963 refer to the calendary car and probably contain some ore needing concentration. (b) Exports. (c) Imports into the United States. (d) Not available.

world production, prices, and imports, see Anon. Mineral Ind. 1935, 44, 58; cf. "Imperial Institute Monograph on Mineral Resources, Chromium Ore, new edition in preparation.

Pure Chromium,-Adeoek (J. Iron and Steel Inst. 1927, 115, 369, 435) obtained the metal of n high degree of purity by the electrolysis of n bath containing 300 g. chromic acid and 10 g. sulphuric acid per litre. A yield of 22.6 g. per kw.-hr. of compact metal was deposited on a rotating steel tube which formed the cathode. This chromium contained hydrogen which was removed by melting the product in vacuo, and oxygen which could only be climinated by heating in a current of pure dry hydrogen at 1,500°-1,600°. The final product, although spectroscopienlly pure, was criticised in the discussion of the paper (l.c. 435). The difficulty of the preparation is shown by the fact that the product could not be melted even in nlumina crucibles in vacuo without being contaminated with oxygen.

Many methods depend on the reduction of chromic oxide. Rohn (Z. Metallk. 1924, 16, 275) heated the oxide to 1,500° in n Heraeus tube furnace in a stream of purified dried hydrogen. The best of these methods is probably the Compt. rend. 1895, 121, 822). thermit process of H. Goldschmidt (Annalen,

For a discussion of chromite ore and chromium, | 1898, 301, 19) in which a mixture of chromic oxido and granular aluminium is brought to exothermic reaction and fusion of the whole mass by firing it with a fuse made of barium peroxide and nluminium powder. The metal, 98-99% pure, separates from a layer of slag.

Commercial Chromium is made by one of two methods, the thermit process above described, and by reduction of chromic oxide with the theoretical proportion of silicon, to which some lime is added to form a slag, in an electric 98-99%, and may contain up to 0.9% Fe and not above 0.8% silicon as major impurities. The metal from the thermit process contains n maximum of 0.9% aluminium. Neither pro-

Pyrophoric Chromium has been prepared by heating chromium amalgam, Hg₃Cr and HgCr, in vacuo below 300°. The amalgam is prepared electrolytically or by the action of sodium nmalgam on solutions of chromic salts (Féréo

Physical Properties .- Electrolytic chromium is

a very hard bluish whita metal which takes a Tables for Spectrum Analysis," 1923, A. Hilger, brilliant polish. The hardness is about 9 on London). Mohs scale, or 2,000 by the scratch test which gives 1,950 for case hardened steel (U.S. Bureau index n 2 97, the index of absorption na 4 85. of Mines, Information Circular, IC. 6566, 1932). The presence of occluded hydrogen has been considered to be the cause of part at least of the hardness (Adcock, Ic.). According to Makarieva and Biryukov (Z. Elektrochem 1935, 41, 623) heating the metal to remove hydrogen may even increase the hardness. The face-centred cubic lattice structure of chromium cenfers maximum hardness, the hexagonal and body-centred cubic lattices being less hard and converted into the first named structure on alcoholic solution of calcium nitrate, it will heating. The density of degassed electrolytic rectify an alternating current (Sborgi and chromium at 25°/4° is 7 138±0 003 and at -50°/4° is 7 156±0-001 (Hüttig and Brodkorb, Z. auorg. Chem. 1925, 144, 347) Döring found 7 085 for 98% chromium made by the thermat process (J. pr Chem. 1902, [u], 68, 65). International Critical Tables, p., 456, and also Landolt-Börnatem Tabellen, 1927, Erganz, 2, 104, give the density as 7 1, for later values, e. soid. Erganz II. i, 210, 111, 1, 282, The coefficient of thermal expansion between 20° and 100° is 0 0000067 (Hidnert Phys Rev. 1932, [2], 39, 186).

Very divergent values have been published in the earlier literatura for the melting point of chromium, 1,420°-2,000°, figures which indicate the difficulty of preparing a pure specimen L. Muller (Ann Physik, 1930, [v], 7, 48) deter-mined the cooling curve of electrolytic chromium with a thermo-couple between 2,000° and 1,400° and found the mp 1,805°. F Hoffmann and Tingwaldt (Z. Metalik 1931, 23, 31), employing an optical pyrometer, observed them p of electro-lytic chromium contained in a magnesia crucible in an evacuated resistance furnace, and also the melting temperature at a hole in a bar of sintered chromium powder heated by an electric current The first method gave 1,800°±10° and the second 1,765±10°. Friemsnn and Sancrwald (Z. anorg. Chem. 1931, 203, 73) found 1,015°-1,925° in rocuo and 1,650° in a mixture of H. and N₄, According to Greenwood (Proc. Roy. Soc. A, 1009, 82, 396) the metal boils at about 2,200°, it can be distilled in an electric furnace under ordinary pressure (Tiede and Birnbrauer, Z. anorg. Chem. 1914, 87, 149) but a

carbide distils in a graphite tube. The apecific heat at -252 5° is 0 014 (Dewar, Proc. Roy. Soc. A, 1913, 89, 158), at -190° values near 0 08 were found by several ubservera (inter alia Richards and Jackson, Z. physikal. Chem 1910, 70, 414); average values at 0°, 0 104, and 100°, 0-110, have been recorded (Schimpff, 151d 1910, 71, 257; Schühel, Z. anorg. Chem 1914, 87, 81). At 1,000° Umino found 0 135 (Sc. Rep. Tóhokw, 1926, 15, 597). According to Jaeger and Rosenbohm (Proc. K. Akad Wetensch, Amsterdam, 1934, 37, 489) thermal

H. von Wartenberg measured the refractive and the reflectivity R%, 697 for the yellow mercury has 579 mµ (Verh. Deut. Physikal. Ges 1910, 12. 105). According to Bernoulli (Physikal. Z. 1904, 5, 632), active and passive chromum give different values for reflexion, a result which suggests the presence of a surface film on the passive metal

Chromium forms a thermoelectric coupla with nickel (Hase, Z. Metallk. 1929, 21, 200). Since ehromsum functions only as anode, eg. in an alcoholic solution of calcium nitrate, it will Cappon, Nuovo Cim., 1922, [vi], 23, 303).

Allolropes of Chromium .- The existence of two allotropes of chromium has been inferred from X ray examination of the metal (Bradley and Ollard, Nature, 1926, 117, 122). Three allo-tropes are reported by Sasaki and Sekito (Trans. Electrochem. Soc. 1931, 59, 437; cf. Makarieva and Buyukuw, Ic). The hexagonal allotrope contained in electrolytic chromium is transformed sate one of the two cubic modifications hy heating the deposited metal to 130° for 15 munutes (Wood, Phil. Mag. 1937, [vul, 23, 984).

Passive and Active Chromium,-Chromium is described as active when it dissolves in the halogen hydracids, oxalic, sulphuric, and hydrofluositicic acids, or when the metal becomes charged with hydrogen while used as a esthode A fresh surface exposed by scratching the passive metal as active. Chrominm becomes passive when ammersed in chlorine, or bromine water, natric, chromic and chloric acids, and in solutions of potassium permanganate, ferric chloride or potassium ferricyanide; when exposed to su or oxygen or if employed as an anode provided the voltage exceeds a critical value. In the electrochemical series passive chromium is placed near Pt, whilst active chromium is close to Zn. Muthmann and Fraunberger measured the potentials acquired by a number of samples in KCI solution against a calomel electrode and found +035 volts with active chromium and -1 47 with passiva chromium; these are the extreme values (Sitzungsber, hayr, Akad, 1904, (ul. 34, 201). Local patches of more or less active ur passive metal may exist on tha sams apecimen. The existence of an oxide film on passive chrominm has been much discussed in the literature (e.g. W. J. Mullar, Z. Elektrochem 1931, 37, 328. For protective oxide films on metals, see numerous papers by U. R. Evans) Among the authors who dispute this explanation are Rathert (Z. physikal. Chem. 1914, 88, 567)

and Bowman (Rec. trav. chim. 1924, 43, 1, 399) Muller and Essin (Z. Elektrochem, 1930, 36, 963-972) measured the potential of chromium, deposited electrolytically on copper, in contact of de Gramont (cf. F. Twyman, "Wavelength with the diameter of the amon, the metal

Further experiments by Müller and Schwabe (Z. Elektrochem. 1931, 37, 185-197) on the activation of chromium by cathodic polarisation in acid solutions show that each acid requires a definite activation potential. The negative potential at which activation occurs in normal acid solutions increases in the order, hydroehlorie < hydrofluorie < hydrobromie < sulplurie < perchlorie < orthophosphorie; thus. activation is more easily affected the smaller the anion is. The potentials for electrolytic and thermit chromium are slightly different. When the activation is affected by grinding with an emery wheel there is again a definite activation potential, which depends on the anion of the acid, and the order is the same as that given above. The experiments lead to the following deductions concerning the nature of passivity. Chromium in the passive state is covered by a network of chromie oxide molecules, which are anchored to the units of the chromium space lattice. Acids cannot dissolve the oxide film except by penetrating between the molecules, and the size of the anion is therefore important.

In a later paper (Z. physikal. Chem. 1936, 176. 273) E. Müller states that in M.HCl at 20°, chromium amalgam oscillates periodically between the active and passive states. In M-H₂SO₄ massive chromium is passive, but can be rendered permanently active by weak cathodic polarisation. The experimental results in the paper are contrasted with the author's earlier explanation of passivity (Z. physikal.

Chem. 1932, 159, 68).
Chemical Properties.—The pure metal is not affected by air, oxygen, or chlorine at temperatures up to 300° C. It is not attacked by nitrie acid, aqua regia, ammonia solution, acetic acid, lactic acid, or by molten zinc, tin, or brass. Chromiam is attacked by gaseous hydrogen chloride, and at high temperatures by sulphur vapour and hydrogen sulphide. It dissolves in hydrochloric acid and the rate of evolution of hydrogen varies periodically with the time, a phenomenon which has been studied by several authors (Ostwald, Z. physikal. Chem. 1900, 35, 33, 20; Adler, ibid. 1912, 80, 385; E. S. Hedges and J. E. Myers, J.C.S. 1924, 125, 601; "The Problem of Physico-Chemical Periodicity," London, 1926). Chromium is also dissolved by the other halogen hydracids, and by dilute sulphurie acid, the action being accelerated by heating, or by the presence of impurities in the metal. The metal burns with vivid sparks when heated in the oxyllydrogen

blowpipe forming the sesquioxide, Cr.O₃.

Electrolytic Chromium, Chromium Plating.—
The brilliant appearance, the hardness and resistance to corrosion of electrolytically deposited chromium are available only since the existence of the modern art of chromium plating, a process which was not generally successful before Fink's U.S.P. 1581188 of 1926. Fink employed a bath containing 250 g. CrO3 so volatile that it disappears from a watch-

becomes active towards the first four acids; 350 g. CrO₃, 2-6 g. H₂SO₄, and 10 g. of in phosphoric, perchloric, and nitric acids it H₃BO₃ per litre and obtained hard deposits remains inactive at 100°. with the bath at 57° and a rate of deposition of 0.002 in. per hour (Metal Clean. Finish. 1935, 7, 325, 385, 535). With a bath resembling Fink's (supra), the hardness increased with falling temperature, 38° or over being too high (ibid. 7, 73, 169; Kronsbein, J. Electrodep. Tech. Soc. 1935, 23; Cymboliste, Compt. rend. 1937, 204, 1654). Poor results are obtained with baths of chromic salts (Applied Chemistry Reports, 1933, 18, 363). Specifications for shroming certains are discussed by Francisch chromium coatings are discussed by Francis-Carter, J. Electrodep. Tech. Soc. 1935, 10, 69. Explanations of the process of electrolysis of the three types of chromium compounds Cr++, the three types of enromatin compounds of the three types of enromatin compounds of the three types of enromatin compounds of the three types of the types of types of the types of types of the types of types of the Wright (Chem. and Ind. 1930, 473) distinguishes two types of chromium plating: (1) on a preliminary coating of nickel or copper for preventing rusting of automobile and machinery parts, taps, cutlery, and scientific instruments; (2) direct plating on tools, pressure rollers, and plates where a hard surface is required (cf. Gardam, Metal Ind. 1930, 36, 245, 279; Dobbs, ibid. 36, 435). The direct method is much less important (cf. Vignos, B, 1937, 798). Disadvantages of the chromic acid bath and suggested improvements are discussed by Cutlibertson (Applied Chemistry Reports, 1937. 22, 413; heavy Ni and Cr plating in England, Fescol process, by Bonilla, Trans. Electrochem. Soc. 1936, 71, Preprint 2, 9). Among special works, see "Chromium Plating," by Bauer, Arndt, and Krause, transl. E. W. Park, E. Arnold, London, 1935; B. Freeman and F. G. "Electroplating with Chromium, Copper, and Nickel," Prentice Hall, New York, 1930; R. Schneidewind, "Chromium Plating, University of Michigan, 1928.

Chromium Hydride. — Electrolytic chromium may contain 250 times its volume of hydrogen (Carveth and Curry, J. Physical Chem. 1905, 9, 353). The gas contained in the metal burns off without further heating after the specimen is brought into contact with a flame (Hüttig and Brodkorb, Z. anorg. Chem. 1925, 144, 341). When an ethereal solution of chronic chloride and phenyl-magnesium bromide is shaken with hydrogen gas, a black precipitate is formed of the composition CrH3 (Weichselfelder and Thiede, Annalen, 1926, 447, 75).

Chromlum Carbonyls.—The hexacarbonyl is formed when carbon monoxide acts on Grignard reagents (e.g. C6H3 MgBr) and ehromic chloride in ether-benzene solution (Job and Cassal, Bull. Soc. chim. 1927. [iv], 41, 1041). After hydrolysis the product is steam-distilled and Cr(CO), is obtained by vacuum sublimation as coleurless, strongly refracting, orthorhombic crystals, isomorphous with Mo(CO)₆ and W(CO)₆, and morphous with morphous soluble in inert organic solvents. It is and 2.5 g. SO₄ ion per litre with a current glass in a few hours. It melts in a scaled density of 34 to 144 amp, per sq. ft. of eathode surface. More recently Piersol has used slowly above 120°, yielding a mirror of pure Cr

(Hieber and Romberg, Z. anorg. Chem. 1935, [LTInd. Chimica, 1932, 7, 1001), a solution of 201 211 When heated with pyridne in a chromic sulphate at 32.5° contains 34.5% of

Cr(CO),(C,H,N),

(Hieber and Mühlbauer, 151d. 221, 337). For organo-metallic compounds of ehromium, see J. Newton Friend, "Inorganic Chemistry," vol. xi, pt. 4, C. Griffin & Co., London, 1937; E. Krause and A. von Grosse, "Chemie der metall-organischen Verbindungen," Berhn, 1937.

Chromium Phosphate, prepared by mixing cold solutions of chrome alum and of disodrum hydrogen phosphate is an amorphous viulet precipitate which left in contact with the solution becomes erystalline and has the com position CrPO,6H,O. On boiling with water it is converted into the green crystalline hydrate CrPO,4H,O. A green amorphous hydrate of this composition is obtained by precipitating a hot chrome alum colution with excess of Na, HPO4. A dihydrate may be prepared by heating the hexahydrate in air, but at low redness an anbydrous black salt and at a higher temperature a green basic salt is obtained (Joseph and Rae, J.C.S. 1917, 111, 1, 196) Green pigments, known as rert Arnaudon and rert Pleasy, have been prepared by redncing a solution of potassium dichromate and aoding a solution of an acid phosphate (cf. Carnot, Compt rend 1882, 94, 1313, Chrome Greens, this vol. p. 107d)

Chromium Sulphate, Cr2 (SO4)3, a crystal-line red powder obtained by heating the hydrated salt above 325° in recue (Krause, Querengisser, and Weyer, infra! It is insoluble in water (Friedrich and Bliekle, Metallurgie, in water (Friedrich and Differie, accessing); 1910, 7, 323), as is the acid salt, Cr.H. (SO₄), (Caley and Burford, Ind. Eng. Chem. [Anal], 1936, 8, 63). Series of violet and of green crystalline bydrates are known and also of green amorphous hydrates. Krauss, Querenglaser, and Weyer (Z. anorg Chem. 1929, 179, 413) quoto from the literature violet hydrated sulphates crystallising with 18, 17, 16, 15, 14, 12, 6, 5, 4 mols. of water, and green hydrated aulphates crystallising with 11, 8.7.6, 5 mols. Their paper describes the prepara-tion of voict hydrates with 18.9, and 3 H₂O, also of the green crystalline hydrato with 12H₂O, and the green amorphous hydrate with 15H2O. The solution of this last salt is precipitated by BaCl, on boiling, but not in the cold. Cold solutions of the violet hydrates are completely precipitated by BaCl. The ordinary violet hydrate with 18H₂O is obtained by crystallising, or by adding alcohol to a solution of chromic hydrate in the calculated quantity of sulphurie acid. Its crystal water has also been reported as 17 mols. H₂O (inter also Weroer and Guber, Ber. 1901, 34, 1592; Montemartini and Vernazus, UInd. Chimica, 1933, 8, 445). The violet solution begins to turn green at 65° and the change is complete on boiling. Alcohol

221, 3-11. Meen beated with pyrindne in a caromic supparts at 3.2° contains 34.5% of sealed tube under suitable conditions it pieds the violet form, between the concentrations bright red Cr(CO)₁(C₂H₁N)₁, sellowsh red Cr₁(CO)₁(C₂H₂N), and pellow 23 days from either side Although byirdytus no doubt occurs in the violet solution (Denham. Z. anorg. Chem. 1908, 57, 361), the change of colour on boiling is probably due to the forms. tion of green complex ions. These have been variously formulated to indicate the masked -SO, groups and the combined water. Zaldes (J. Gen. Chem. Russ. 1936, 6, 1325; A.I. 1937) writes

> (Cr(OH,),1,(SO,),+H,O → [Cr(H,O),(SO,)]+H+[Cr(H,O),OH]" +50

Rasic complexes in chromic sulphate solutions are considered to be effective in chrome tanning (J. A. Wilson, J. Amer. Leather Chem Assoc 1936, 31, 165; cf. Stiasny et al. Chem Assoc 1990, 31, 105; 6. Suasny et at., Colleguam, 1928, 49, 72). The green crystalline sulphate, Cr, (SO₄), 12H₂O, is prepared by passing SO₂ into a concentrated CrO₂ solution at 6°-5° and crystallising in vacuo (Krauss et at., Lc). In a series of papers in the Comptes rendus from 1905 to 1907, Colson described the preparation of green bydrated sulphates with respectively 3, 2, and 1 mesked SO, groups, the two latter salts being successive stages in the hydrolysis of the first, [Cr.(SO,1)] Recours (Ann Chim 1895, [vu], 4, 511) prepared the mono, di, and trichromosulphuric acids by heating a green crystalime bydrate, Cr₂(SO₄)₂,8H₂O, with 1, 2, or 3 mols, of sulphurio acid to 110°-120°. The acids are resepctively di, tetra, and herabasic, corresponding to the proposed formula [Cr₂(SO₄)₃]H₄. They dissolve to deep green solutions in water which if freshly prepared do not give the reactions of Cr or SO, ions, According to Küntzel et al. chrominm sulphate complexes are contained in a boiled and cooled chromic sulphate solution, Masking of chromic salts occurs on adding a nentral salt containing an anion common to both (Collegium, 1035, 484). Montemartini and Vernazza (I c. 8, 445) prepared a saturated solution of chromic sulphate, Cr. (SO4), 17H,O, at 18° containing 84 8% of this salt, and state that the equilibrium violet green was not affected by addition of alkali sulphate, the percentage of violet salt in this solution was

51 68 Chrome Alums .- The double sulphates of chromium and potassium or ammonium, M,SO4.Cr, (SO4),24H,O, are important chemicals; they crystallise in large violet octahedra and are very soluble in water. The corresponding sodium salt does not crystallise so readily. The ammonium salt is made by mixing molecular proportions of the constituent eulphates, potassium chrome alum is a common by-product from oxidations with potassium dichromate, although on the large acale, eq the change is complete on boiling. Alcohol authracene → authraquinone, the chromic acid now precipitates a green basic salt (Venable and solution is regenerated by electrolytic oxidation Miller J. Amer. Chem. Soc. 1898, 20, 484), of the chromo sulphate solution. Potassum The green solution becomes violet on standing, chrome alum melts at 89°, turning green According to Montemartini and Vernazza (Tiden, J.C.s. 1884, 45, 409); in solution the Vernazzi (l.c. 8, 445) state that the saturated solution at 18° contains 28.2% of

K2SO4.Cr(SO4)2.24H,O,

of which 51.75% exists as a violet salt; the saturated solution of the sodium ehrome alum contains 94.8% of the salt with $24H_2O$, of which 51-7% exists as a violet salt. Chrome alum is used in paper-making, photography, dyeing, printing, and tanning. Immersion in a bath of chrome alum hardens gelatin films and renders them insoluble, an effect attributed to the fixation of chromic hydratc.

Pure crystallisable violet ehrome alum may be prepared by the electrolysis of a solution of potassium dichromate mixed with sulphurie acid. The liquor is circulated in the cathode compartment which is separated from the anode by a porous vessel. The cathode is preferably of carbon. Dilute sulphuric acid is circulated in the anode compartment and the anode may be eopper with copper turnings to utilise the SO₄ ions (Chaumat, F.P. 459677, G.P. 265170).

Chrome alum is manufactured from ferrochromium by Chemisehe Fabrik in Billwarder, vorm. Hell and Sthamer A.G., and Hasenclever, by dissolving the alloy in sulphuric or hydroehloric acid and adding to the solution sufficient alkali to precipitate only the chromium present; the precipitate is converted into chromic sulphate and chrome alum in the usual way (B.P. 187232). In a similar process the same patentees separate about half the iron by erystallisation as ferrous sulpliate and centrifuging. The liquor, from 1,000 kg. of ferro-chromium and 3,000 kg. of sulphuric acid, is next treated with potassium sulphate, 10 kg. of petassium dichromate, and 3 kg. of sulphuric neid, while sulphur dioxide is passed into the liquor, ehrome alum then erystallises (B.P. 187231). The erystallisation of chrome alum is accelerated if the potassium sulphate is added to a solution of chromic sulphate previously acidified with sulphuric acid and heated to 30°-45°, and the solution then cooled rapidly while stirring (I.G. Farbonind, A.G., B.P. 260885). For use as a mordant, a solution of chrome alum is readily prepared by adding sugar or glucose to a hot solution of potassium dichromate and sulphuric acid. Another process depends on the insolubility of a basic chromic sulphate. Aqueous chromic sulphate containing ferrous sulphate produced by dissolving ferrocliromium in sulphuric acid is treated while hot with hydrated lime to neutralise the free acid and filtered from CaSO4. filtrate diluted to 4-5 times its volume with hot water deposits basic chromic sulphate which can be economically converted into Cr2(SO4)3 and chrome alum (W. Hene, B.P. 418714).

A violet chrome alum solution requires the normal amount of NaOH for complete precipitation, but after boiling, only & of the theoretical amount. Non-ionised SO, in the complex varies from half the total SO, in a (Atkin J. Chollet. Soc.

change occurs at about 78°. Montemartini and [Trades' Chem. 1934, 18, 356; Ch. Abs. 1934, 6078). Chrome tanning, as affected by the formation of complexes and by the solubility of basic chromic sulphates, is discussed, by the same authors (Cuir tech. 1934, 23, 142; cf. Küntzel, Kinzer, and Stiasny, Collegium, 1934, 213). The $p_{\rm H}$ of solutions of basic chromic sulphate and the absorption of ${\rm Cr_2O_3}$ therefrom during tanning has been studied by Britton ("Hydrogen Ions," 1929). Basic chromo alum solutions are used in the one-bath process of chrome tanning. Following earlier authors, Lumière and Seyewitz (Bull. Soc. chim. 1903, [iii], 29, 1077) found that Recoura's green basic sulphate, 2Cr₂O₃,5SO₃, hardens gelatine more efficiently than chromo alum.

Chromium Chlorate, Cr2(ClO3)6.-Solutions of this powerful oxidising agent have been prepared in calieo printing works by mixing solutions of chrome alum and potassium or barium chlorate. It has been suggested for the production of ungreenable aniline blacks, but is considered to be liable to tender the fibre. The solution has a strong smell of chlorine and gives off oxides of chlorine when heated. It has been proposed as a mordant for catechu browns (cf. Kneeht and Fothergill, "Textile Printing," 1st ed. 1912, 201; 2nd. ed. 1924, 211, C. Griffin & Co., London).

Chromium Dioxide, CrO2, and intermediate oxides, $xCr_2O_3yCrO_3$. Manchot and Kraus (Ber. 1906, 39, 3512) heated blue chromic chloride hydrate, precipitated from chromic chloride solution by ammonia, to 330° in a current of oxygen, and obtained a black hygroseopie powder which on heating to a higher temperature glowed and evolved a volume of oxygen corresponding to $4\text{CrO}_2 = 2\text{Cr}_2\text{O}_3 + \text{O}_2$. L. Blanc (Ann. Chim. 1926, [X], 6, 182) precipitated brown $2\text{Cr}_2\text{O}_3$. CrO₃= Cr_6O_9 by adding a solution of a chromic salt to a solution of potassium chromate, but not if the order of mixing is reversed. A table of these intermediate oxides has been compiled by Blanc (l.e., 199).

Chromlum Chromate, CrO3, Cr2O3, 7H2O. -Jovitschitsch (Helv. Chim. Acta, 1920, 3, 40) states that when a solution of chromic oxide in HNO₃ is evaporated until excess of acid is removed and the residue dissolved in water and treated with ammonia a brown substance is precipitated which is black when air-dry. Analyses give the formula $Cr_2(OH)_8$, H_2O or CrO_3 , Cr_2O_3 , $7H_2O$. It loses $1H_2O$ over H_2SO_4 , $2H_2O$ at 105° , and $4H_2O$ at 205° . By dissolving Cr₂O₃ in HNO₃ and evaporating and heating to 290°, a black product is obtained believed to be CrO₃·Cr₂O₃ (cf. Jovitsehitseh, Monatsh. 1912, 33, 9; 1913, 34, 225). For other preparations, see under chromic oxide, and the decomposition of CrO₃ by heating.

Chromic Anhydride, Chromic Acid, CrO₃. -This oxide is usually prepared by mixing excess of concentrated sulphuric acid with a saturated solution of potassium or sodium dichromate or by digesting barium chromate with complex varies from half the total SO, in a dilute sulphuric acid and evaporating the boiled 1% solution to two-thirds the total in a filtrate. The crystals which separate contain boiled solution of 150 g./l. This latter solution sulphuric acid if made by the first process and contains the complex [Cr4(OH)2(SO4)412H2O] either barium dichromate or sulphuric acid if Leather made by the second, and must be purified by

453) mixed 10 volumes of rold saturated potassium dichromate solution with 12-15 volumes of cone, sulphuric acid, and pollected the separated crystals on a filter of broken glass transferred them to porous plates and recrystallised the crude product from a little water. Bunsen (Annalen, 1868, 148, 290) mixed 5 parts of sulphuric acid with 11 parts of a solution rontaming 1 part K,Cr,O, after 24 hours the CrO, crystals which separated were filtered off by suction, washed with nitre acid, and heated in a current of dry air at 60°-80° Zethow (Pocc Ann. 1871, 143, 468) states that Zethow (Pogg Ann. 1871, 143, 469) states that he nitre suid should be not less than 1-16 spgr. An anonymous author (J Chem. Eng. China, 1305, 3, 39) treats K₁Cr₂O₂ with 88% intro acud and separates the CrO₃ from KNO₃ by fractional crystallesshor. The yelds 199% of CrO₃ of 98% purity, suitable for chromusin juding baths (f. Lee and Kung. B. 1937, 901)

Metals Protection Corporation (B P. 307061) wash the crude CrO, with sulphurie acid to remove NaHSO, or remove the greater part of adhering mother liquor by heating, and pre cipitate residual sulphuric acid by adding Ba(OH), BaCO, or BaCrO, Bolley (Annalen, 1845, 56, 113) had already pointed of the composition H,SO4+H,O Several

methods start from calcium chromate.

hgof, Korzuna, and Astroy a bouleslesum chromate until it is converted into the difficultly soluble 2CaCrO, H,O, which is nitered off, uashed, and decomposed with sulphure aced, the filtrate, and decomposed with sulphure aced, the filtrate, and the CaSO, is treated with sulphure aced, and CrO, crystallised and centrifuged (Khim At ann From. 1934, 1, 38, cf. Rakorski, J. Phys. Chem. Russ. 1928, 60, 13) Vetter, assr. to Natural Products Refining Co. (U.S. P. 2034256).

describes the rouditions for preparing CrO₃ of high purity by the action of H₂SO₄ on Na₂Cr₂O₅. Polyanski treats a solution of calcium chromate (2 pt) with H₂SO₄ (1 pt d 184) in successive portions, heating on the water both and removing CaSO, between cith addition, until no more is precipitated; hadly CrO₃ is crystallised and purified by Bunsen's method (r. supra, Trans. Inst. Pure Chem Reigents, Sci Tech. Dept B.SSR 1929, No 300, 143) Tanahashi (Japan P 99139, 1933) treats sodium dichromate with 4-5 times its equivalent of sulphume seid of 55°-58° lie, sodmni hydrogen sulphate dissolves while CrO, is precipitated in fine crystals, which are

envisibleston after addition of the required sodium chromate solution is rontinuous, quantities of citeth return chromator suphanie fod into the lover part of the cell, while NaOH and. Commercial "pure" chromic and has and ErO, solutions overflow separately from been known to contain sulphate (Adeed, Am of the upper compartments. Angelini and Pare Iron and Steel Inst. 1927, 115, 369). In one baseo (F.P. 770734) heat chromite mused with carly method Wanngton (Flb. Mag. 1842, 30), asswitss or peat to 400°, chlorine is then passed sawdust or peat to 400°, chlorine is then passed over the porous material. It was already Lnown (Moissan, Compt. rend. 1880, 90, 1337) that chromyl chloride, CrO,Cl, is formed in a similar reaction. With water this gives CrO. and hydrochloric acid. Udy (assr. to Electro Metalling Co., U.S.P. 1878918) claims an electrolytic process in which an anode of ferrochrome dissolves in aulphuric acid. d 1 38-1-52 contained in the anode compartment. CrO.

is en stallised from the acid. Chromic anhydride (often termed thromic acid) is a strongly acid substance crystallising in scarlet rhombie bipyramids. Its formula is of any compound (Buchner and Prus, Z. physikal Chem 1913, 81, 113). The crystals have a specific gravity variously stated, with 2 737/14° and 2 819/20° as limits. The highest value rerorded for the melting point is 198°, (Jaeger and Germs, Z anorg Chem. 1921, 119, 145) When heated to 125° under 16 mm pressure it subhmes in rod needles (Arctowski. soid 1895, 9, 29). At a temperature above but near its melting point it decomposes into chromie oxide and oxygen Above 250° out that CrO, is least soluble in sulphuric acid it is said to yield a residue of the oxide Cr.O. (Cr.O. 3CrO.) which may be purified by boiling with water According to Simon and Schmidt (Z anorg Chem. 1924, 152, 191) this oxide is formed between 370° and 450°. Its Kirkexistence is supported by the experiments of Honda and Sone (Sci. Rep. Tohokw, 1914, 3, 223). A saturated solution of chromic acid at 15° contains 62 4 g, CrO₃ in 100 g, solution The nature of the ions in dilute solutions of chromie ecid has been the subject of many investigations, but the results are in conflict Britton (JCS, 1924, 125, 1572) from electrometrie titration reneluded that in dilute solution the sons are H+ and HCrO,-, Spitalsky's results are in agreement (Z anorg, Chem, 1907, 54, 265) The dissociation to the second stage HCrO₄ = ⇒ H++CrO₄ = is extremely small Other authors, including Ostwald, have argued that the aqueous solution contains Cr.O,-

Chromic anhydride is a powerful oxidiving ageot. Warm, anhydrous alcohol inflames when dropped upon at, and even dry ammonia gas is oxidised with production of water and nitrogen. Addition of evalue acid is said to increase its oxidising power. The solution in water, sulre-re-tailised from dilute suprairie acid and ordinary agent in organie work. In martiel with a saturated solution of CrO, to ordinary agent in organie work. In martinory solphure acid, J. W. Boss (C.S.) these availations precautions must be taken to remove substance and the solution of the reaction. When layer in a cell of which the upper half is divided to be oudised is usually dissolved in the same by an imperincable partition into an anode solvent, and the exidering solution slowly added, and criticals connectionat containing respectively as up ar later element and solution and solution and solution and solution and solution call of d 12. The ing agent, his the advantage that the end of

green colour.

It has the property of producing a permanent yellow colour on silk or wool, but not on cotton. The chromium in such material may be converted into lead chromate upon the fibre or further dyed by logwood, etc. Chromic anhydride usually contains sulphuric acid and

sulphates or nitrates.

Chromates.—All chromates are prepared directly or indirectly by the action of exidising on ferrochromo or on FeO, Cr2O3 (q.v.). Both the normal and acid chromates are of importance industrially. The normal chromates of sodium and potassium, M₂CrO₄, are readily converted into the di-chromates, M₂Cr₂O₇, by the addition of sulphuric acid to their solutions, which then change from yellow to orange. A basic chromate, orange chrome, PbO·PbCrO4, is also manufactured in connection with chrome yellow, PbCrO₄.

A great number of processes have been proposed for the conversion of Cr_2O_3 in chromite into CrO_3 ; the oxidising agent is now atmospherie oxygen acting on an alkalino fusion

of the ore.

In former times the ore was mixed with about two-fifths of its weight of potassium nitrate and ignited on the bed of a reverberatory furnace, the oxygen required for the exidation of sesquioxido into the anhydride and the potassium for its conversion into potassium chromato being both provided by the nitrate. A great saving was introduced by the substitution of atmospheric oxygen as the oxidising agent in place of the nitre, the potassium being supplied by potassium earbonate or, as proposed by Watt, hy potassium sulphate (cf. Gibbs, assr. to National Electrolytic Co., U.S.P. 901436, 1908).

In the modorn process a mixture of chromite, soda ash, and limo is heated in a controlled supply of air to a high temperature in a rever-

beratory or in a rotating furnace.

Sofianopoulos (J.S.C.I. 1930, 49, 279T) recommends a furnace charge of a finely powdered (60 mesh) mixture of 340 lb. of chromite (53.6% Cr₂O₃), 270 lb. soda ash (92%), 300 lb. lime (85% alkalinity), and a reaction temperature of 1,400°F. Under these conditions the recovery of soluble sodium chromates was 91.5% of theory, and the method of working described in the original avoids as much as possible the loss due to production of calcium chromate, which in practice was found to be very slowly soluble in boiling water.

In another method of working these difficulties are overcome; fusion of the charge is avoided and the ealcium chromate produced is boiled with sodium carbonate in vats fitted with stirrers. The filtrate is concentrated in vacuo, and a slight excess of sulphuric acid added, when the greater part of the sodium sulphate separates, the remainder is removed during further concentration, and finally sodium debromate crystallises

covered if the sedium chromate is converted into form of hexagonal lamelle.

the reaction is frequently indicated by a pure | dioxide under pressure so long as sodium bicarbonate is precipitated. This salt then returns to the process and the conversion of chromate into dichromate is completed by the addition of the required amount of sulphuric acid. The less soluble potassium salt is obtained by adding potassium chloride to the solution of Na2Cr2O2. Fusion of chromite with caustic soda has been suggested (G.P. 151135, 163814, and 171089). A direct production of sodium dicbromate has been claimed by Bozel-Maletra Soc. industr. de prod. chim. (G.P. 625568), in which the chromite ore is heated in a rotary furnace with soda-ash in a current of air. In a wet process owned by the same Company (J. E. Demant assr.) the chromite is treated with oxygen gas in an aqueous medium under pressure at temperatures above 100° in presence of Na2SO4 or other salt yielding active eations (U.S.P. 2012061), or ferrocbromium, and NaOH, KOH, or K₃PO₄ are the starting materials (U.S.P. 2012062, B.P. 416624). The anodic oxidation of chromic salts to chromates has been studied by Gross and Hickling (J.C.S. 1937, 325), who explain the results by assuming a primary formation of H_2O_2 at the anode. Spent chromic acid baths from exidation processes are regenerated by electrolytic oxidation in the anode compartment of a divided cell by a method invented by the I.G. Farbenind. A.-G. The same patentees manufacture potassium dichromato by heating a mixture of chromic oxide and potassium hydroxide to 300°-380° in a revolving pan in presence of air or oxygen (B.P. 416744). According to another patent of the same origin, sodium dichromate is manufactured from aqueous sodium chromate containing 475 g./l. of CrO₃ by treatment at 60°-80° with CO2 under pressure, and cooling without releasing the pressure. The conversion of mono- into dichromato is thus 90%, while NaHCO₃ separates and is removed. The filtrato is concentrated to 800-820 g./l. of CrO3 and again treated with CO2 to complete the conversion into Na2Cr2O7 (1.G. Farbenind. A.-G., B.P. 424821).

The dry method of chromate production has been investigated with the following results:

Calcium and chromic oxides begin to interact in the presence of air to form calcium chromate at 650°; at 700° a 95% yield of chromate is obtained with mixtures containing 2 equivalents of CaO to 1 of chromic oxide and a 60% yield when the ratio is 1:1. Pure calcium cbromate does not begin to decompose until 1,000° is reached. Sodium carbonate and chromic oxide readily react to form chromate at a temperature below 660°; the salt melts at 800° and remains undecomposed after prolonged heating at 1,000° (M. R. Nayar, H. E. Watson, and J. J. Sudborough, J. Indian Inst. Sci. 1924, 7, 53).

ALLOYS .- Various alloys of chromium have been prepared, chiefly by reduction in the electric furnace, by the "thermit" method, or by the usual mixture of molten metals,

Molten zine dissolves but little chromium; The sodium carbonate may be partly re- a hard and brittle alloy has been obtained in the Aluminium and dichromate by passing into the solution carbon chromium mixtures, containing between 5 and

55% chromium, separate into two hquid layers, and probably contain a compound, CraAl. The alloys with a low percentage of chromium are hrittle, one containing 13% chromium can be

powdered in a mortar. Alloys with antimony are hrittle, and com-pounds represented by ShCr and Sh₂Cr and Homogeneous alloys, with 30% Cr, when cooled known. Chromium alloys with opper with separate into two acts of crystals, Lewkonja, Z difficulty, but an alloy can be obtained by attrring in copper oxide into a molten alloy of

metals are miscible only to a small extent. Silver and chromium, although partially miscible in the liquid state, form no solid solution Molten eadmium does not dissolve any chromium. Molten mixtures of lead and chromium, containing more than 27% lead, separate into this mixture and pure lead.

Platinum-chromium,-V A, Nemilov (Z. anorg, Chem 1934, 218, 33) finds that the Brinell hardness-composition curve of Cr.Pt alloys shows well-marked minima corresponding with exist. The temperature of polymorphic and compounds Cr. Pt and CrPt Examination is maximal for 75% Cr. falling of the reserve attructure confirms the existence with a higher chromium content. The temperature of the latter Alloys quenched fcom a high temperature show only reuzed crystals, from which CrPt separates on annealing annealed specimens there is a continuous series of roived crystals between 0 and 25 atomic % and another between 65 and 100 atomic % Pt. The electrical resistance rises very steeply with increasing Cr content up to 13 atomic % Cr, hey ond which the alloys are no longer ductile Alloys containing 4-5% Cr have about five times the resistance of Pt, and a higher reciting point H₃SO₄, HCl, and HNO₃, dilute or conc, show no action either at room temperathan 14 atomic %.

Chromium-nickel.-G Voss (Z. anorg Chem 1908, 57, 34) finds that the system CrNi is homogeneous in the liquid phase with a series of mixed crystals in the solid state, the curve showing a deep minimum at 1,290° and

40-42% nickel. In the system chromium nickel, Matennaga (A, 1930, 680) states the components are mucible in all proportions in the liquid phase. in the solid phase the cutectie m p 1,346° contains 49% Ni. The best workable acid reasting

alloy contains 15-35% Cr. X ray investigation of the nickel chromium system by E. R. Jette, V H. Nordstrom, B. Queneau, and F. Foote (Amer Inst. Mm. Met. Eng, Inst. Mct. Div., Tech Pub, 1934, No. 522) shows that below 1,150° there are two terminal solid solutions with an intervening two phase area. On the Cr aide the solubility of N1 in Cr is low, but increases rapidly at over 900°. The solubility of Cr in N1 increases uniformly and rapidly with rise in temperature, that at 1,153° heing nearly 53 wt.% Cr, which is beyond the cutectic point as determined by thermal analysis. The only structures observed were body-centred Cr, face centred Ni, and occasionally rhombohedral Cr₂O₂ (cf. Nishigori and liamasumi Sci. Rep. Tohoku Univ. 1929, 18, 491).

nickel are non-magnetic. Cohalt-chromium.-Cobalt and chromium are miscible in all proportions, both in the hould and the solid states, and yield a mixture of minimum melting point 1,320° with 47% Cr. Homogeneous alloys, with 30% Cr, when cooled

Nickel-chromium alloys with less than 90%

anorg Chem 1908, 59, 325.) Weser and Haschimoto (Chem. Zentr. 1930. aluminium and chromium (Moissan). The hound I, 3482), in reconstructing the equilibrium dragram for the system cobalt-chromium, state that on account of the amail velocity of diffusion of the components complete equilibrium is attamed only with difficulty. Chromium has mp 1.765°+10°. Cobalt and chromium are miscible in the fused state in all proportions The eutectic is at 1,408° and 42% Cr. Up to 38% Cr the mixed crystals are face-centred cohic, whilst above 48% Cr they are body. centred. The compounds Co.Cr, and CoCr (tetragonal, with S mols in the unit cell) probably ture of magnetic transformation falls almost proportionally to the chromium content. The resistance to corresion by acid is considerable, being greatest for nitric and least for hydrochloric acid (for the industrially important CrCo alloys, see this vol. p. 217b, Stellite

Alloys), Copper-chromlum, - In the system of copper cheomium there is only partial reiscibility in the liquid state with a cutectio near the copper end (1-5% Cr, 1,076°), the limit within which no mixture of the two liquids is formed helog 37-93% chromium, above 1,470°. In the solid state ture or at their boiling points, on poished only heterogeneous mixtures of two crystal surfaces of alloys with a Cr content not greater apecies are obtained, chromium and the eutocic rich in copper. In the ternary system Cr-Ni-Cu the heterogeneity of the Cu-Cr system is 55% removed by addition of NI (Siedschlag, Z anorg Chem, 1923, 131, 173)

Chromium-molyhdenum (Siedschlag, ibid 131, 131-3| olyhdenum dissols ea readily in chromshowed a cutectic at 1,460°-22.7% Mo. The curve liquid is homogeneous, but no mixed crystals are formed below 227% Mo, the solid consisting of chromium crystals and the entection mixture Above this proportion the solid conaista of the entectic and crystals of molybdenum

containing 2 5% chrommim. Carhon-chromlum .- These alloys fall into two groups . (I) Those containing 8 5% carbon completely soluble in hot 24% hydrochloric acid; (2) those containing more than 85% carbon, partly soluble in the same acid, obtained by melting chromium (prepared by the aluminium thermse process) in a crucible of pure carbon at temperature varying from 1,840° to 2,567° for 15 minutes and stirring with a carbon rod The alloys of the first group leave a residue of practically pure Cr. C, which forms alvery crystals, d. 6915, mp. 1,665°. No trace of Cr. C, draenbed by Mossan, was found. The alloya containing more than \$ 5% carbon contain no free chromium, only carbide and graphite. The carbon content of the asturated

alloy at 1,840° is 12.42% earbon; at 2,233° 14.03%, and at 2,507° 23.12%. With 24% HCl an alloy dissolves, indicating Cr4C2. The insoluble residue contains graphite and Cr₃C₂. This is resistant to all acids and is darker than Cr_5C_2 , d. 6.083, m.p. 1,890±10°. It is decomposed by chlorine at a red heat, forming the trichloride and earbon. The chemical composition of the alloys rich in carbon is unaffected by the rate of cooling, but larger erystals are developed when the cooling is at 2,570°/8-14 mm. pressure, the vapour being pure ehromium (Ruff and Foehr, Z. anorg. Chem. 1918, 104, 27-46). Maurer and Nienhaus (Stahl u. Eisen, 1928, 48, 996), from collected analyses, show that the earbides of chromium

are Cr_3C_2 , Cr_4C_2 , Cr_5C_2 .

Kraiczek and Sauerwald (Z. anorg. Cbem. 185, 193) investigated the system ehromium-earbon between 0-13.3% earbon by means of heating and cooling ourves and examination of micro-structure and density and observed that in the interval between 8.32-9.9% earbon mixed crystals were formed and a transformation occurs at 1,465°. The evidence indicates the existence of Cr_5C_2 and Cr_3C_2 , but not Cr_4C or Cr_4C_2 . Westgren and Phragmin third 187 4011 2144 214 men (ibid. 187, 401) state that the chromiumchromium carbide cutectic has 3.4% carbon, not 4.5%. A cutectic carbide of the formula Cr, C3

is indicated in addition to Cr₄C,Cr₅C₂.

Schenck, Kurzen, and Wesselkoek (Z. anorg. Chem. 1932, 203, 159), in studying the synthesis of earbides from the metals and CH4, find that CH4 reacts with Cr at 600°-800°, forming Cr₅C₂ and a C-rich mixed-crystal phase containing 11.5% C in equilibrium with the Cr₅C₂ phase. The C content of the mixed-crystal phase can be increased to 12.5% by further treatment with CH4; this phase

probably contains Cr₃C₂.

By means of the equilibrium diagram of the chromium-earbon system, Hatsuta (J. Study Met. 1931, 8, 81-88) finds four earbide phases, ϵ , η , ζ , and κ , corresponding respectively with Cr_4C (enbie), Cr_7C_3 (trigonal), Cr_3C_2 (orthorhombic), and (?) CrC. A cutectic $(\alpha+\epsilon)$ lies at 1.485°, 3.79% C. The ϵ and η phases are peritectically formed thus: $\eta + \text{melt} \rightarrow \epsilon (1,530^\circ)$, $\zeta + \text{melt} \rightarrow \eta$ (above 1,600°). The ζ phase probably has the maximum melting-point on the liquidus eurve, and forms a euteetie with the κ phase; a transformation takes place at 1,505° in this phase.

lron-Chromium .- With iron the alloys of chromium are of great interest. The system Cr-Fe on solidification exhibits an unbroken series of mixed crystals. aFe-Cr and &Fe-Cr show complete miscibility, while yFe (stable between 906°-1,400°) forms mixed crystals with Cr up to 14% (Bain, Trans. Amer. Soc. Steel Treat. 1926, 9, 9; Oberhoffer and Esser. Steel freat. 1920, 9, 9; Obermonet and Seel, Stahl n. Eisen, 1927, 47, 2021; Maurer and Nienhaus, ibid. 1928, 48, 996; Wever, ibid. 1929, 49, 839; Chem. Zentr. 1931, ii, 2924; Bain and Griffiths, Trans. Amer. Inst. Min. Met. Eng. 1927, 75, 166). The presence of chromium in iron or steel produces a much finer texture,

greater smoothness of fracture. Chromium lowers the Ar, point, eventually causing its disappearance, whilst it raises the Ar point of the steel, the proportion of chromium required being less the bigher the earbon, e.g. with 0.2% carbon, 5% chromium is required. Henco chromium steels are self-hardening (i.e. air-hardening). Steels containing 15% chromium have been used for tools and have to be east to shape. With 0.5% chromium and 0.5% earbon tho steel is used for artesian well bits and jars and retarded. An alloy saturated with earbon boils has no equal in the hardened state. With 1% chromium and high carbon the steel is used in bearing balls, cones, roller bearings, and crusbing machinery. These low chromium steels, when well annealed, can be machined. The principal use of ehromium is as a constituent of other alloy steels especially nickel steels, to which it imparts greater toughness and hardness. As such it is used extensively in automobile parts and in armour plates which are case-hardened and quenched. Chrome steel is also used in the manufacture of special kinds of files. The alloy is usually prepared of the requisite composition by the addition of a definite amount of " ferrochrome," containing from 40 to 85% chromium, to the molten steel.

Stainless steel cutlery containing Cr 11-14% and C 0·3-0·4% was introduced in 1913. For chemical plant stainless steel may contain Cr 8-18% with Ni 8% or Mn 4%. These alloys resist nitric acid, acetic acid, caustic soda, and ammonia. The steel with 14% Cr and 0.15% C will resist hot but not boiling nitric acid. Wires of a chrome steel containing a high percentage of nickel can be fused into glass liko platinum. Laboratory electric furnaces are usually wired with Ni-chrome resistance wire (A. B. Kinzel and W. Crafts, "The Alloys of Iron and Chromium," I, 1937; "Low Chromium Alloys," II (now published), "High Chromium Alloys," McGraw-Hill Book Co. Inc., New York and London, 1937; R. H. Greaves, "Chromium Steels," H.M. Stationery Office, London, 1935). For alloys of chromium, see Priestloy, Ind. Eng. Chem. 1936, 28, 1381; for chromium steel in pressure vessel construc-tion, see Hopkins, ibid. 28, 1386.

Ferrochrome was formerly prepared from rieb chrome iron ores in blast furnaces by the use of coke and hot high-pressure blast, or in erucibles. In 1890 the manufacture was commenced in the electric furnace and has gradually superseded the other methods. In the production of eliromium from eliromite to form ferroehrome, the iron oxide being reduced, reduction of chromic oxide begins about 1,185°C. Thirty parts of pure earbon are theoretically necessary for every 100 parts of iron and chromium reduced. On the basis of the double carbide Fe₃C,Cr₃C₂ being formed by reduction of the eliromite the following reaction occurs:

9FeCr₂O₄+50C $=[2(Fe_3C,3Cr_3C_2)+3Fe]+36CO$

The ferrochrome resulting would contain theoretically 10.4% carbon, 31.4% iron, and 58.2% chromium. The ore has to be finely ground and is then mixed with anthracite greater hardness, tenacity, and elasticity, and coal in the proportion of 37 parts of carbon to

сивомили.

100 parts of are, which forms the above alloy, of chromic hydroxide, chromic anhydride, and and charged into the furnace. The power certain chromotes. consumption in a 750 kilowatt furnace of the Alby carbide type at Koppersen, Norway, was potassium chromete or dichromete with a 30 km, hours per lh, or 0-68 km, year per short reducing egent, etarch, glyrenne, suger, etc. ton for 5% carbon and 65% chromium ferro-chrome Tho percentage of carbon in the ferrocharged unto the furnace, but decarburisation can be effected with on oxide alag of iron or chromium after tapping the elag from the first reduction (Trans Electrochem. Soc. 1913, 176).

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S. Eriksson (Jernkont, Ann. 1931, 118, 530; A. 1935, 1455) confirms the existence of an intermediate a phase in the Fe-Cr system (approximate formula FeCr) (cf. Wever, B. 1932, 469).

The addition of chromium decreases the mognetie properties, but all alloys, to 80% chrominm. are megnetic. The magnetic property of chromium is not increased by cooling in solid hydrogen (Compt, rend. 1901, 150, 687). A earhido of chromium and tungsten of great hardness and of ap gr. 841 has been produced in the electric furnace, and to it the special properties of chrome tungsten steels ore probably due. These latter (as well so chrome molybdenum steels), containing up to 3% chromium end 16%; tungsten, are used for the manufacture of machino tools Cohalt is frequently added to both classes of tool steel Steel containing 12% Cr may be produced by reducing Cr.O. by ferro silicon shove a hath of low-carbon steel (Applied Chem. Rept. 1936, 341)

Basic Oxides,-Two chromium oxides, oxides, Jielding salts with acids in which chromium is present as a divalent or trivalent kation, forming the chromous and the chromic Chromio oxide can also combino with bases to form salts, the chromites, of the type MCrO, in which the chrominm is present in the monovalent amon, CrO2. Chromic anhydride, CrO2, is an acid anhydride forming, with bases, salts of the type M.CrO., the chromates, or M.Cr.O., the dichromates, in which tho or Cr.O., Other oxides have been prepared and ore usually regarded as compounds of the basic and set usually regarded as compounds of the chromie chromate or chromium dioxide Cro. These oxides are described on p. 101,

The asits of chromium are coloured ahodes of violet or green; all solutions of violet chromium salts show similar absorption towards light; the spectrum is due, therefore, to the chromium In solutions of green complex salts the absorption band and limit of complete absorption are both shifted towerds the red end of the

spectrum.

Chromous Oxide, CrO, is most conveniently prepared by the action of dilute nitric ecid on chromium amalgam which dissolves the mercury leaving chromous oxide as a black powder (Dieckmann and Hanf, Z. annry Chem 1914, 86, 301).

Chromic Oxide, Chromium Sesquioxide, Cr.O. -This compound is produced by the oxidation of metallic chromium and by ignition 425), a very pure colour, suitable for colouring

(1) Amorphous: (a) Prepared by heating and washing the residue with hot water. (b) By heating ammonium dichromate, according to a number of authors the product is green Cr.O. This reaction has been studied by Herbard and King (J.C.S. 1938, 955). In vacuo the product was a black powder containing Cr 61.2% (Cr.O. contains 68 42% Cr); on further heating an olivecoloured substance was obtained (Cr 673± 05%), this heated in hydrogen'at 800° for 21 hours yielded hright green chromic oxide (Cr 684%) The thermal decomposition of (NH₄)₄Cr₁O₇ appears to yield e non-stoicheiometric compound (cf. Bell, J.C.S. 1909, 95, 87, who by slow de-(cf. Bett. d.L.o. 1893), 53, 61, nao by sive uc-composition of (NH₄), Cr.O. ohtained a black powder, 3CrO₂·H₄O; Hooton, Proc. Chem. Soc. 1908, 24, 27, found H.Cr.O.; Moles and Gonzalez (Anal. Fis. Quím. 1923, 21, 204) reported the product as CrO₂. Fischbeck and Spingler (Z. anorg. Chem. 1938, 235, 185) confirm Hooton's formula. For criticism of these results, see Harhard and King, I c.). (c) By heating mercurous chromste, preferably with out access of air. (d) Not in the pure state, by heating chromic hydroxido in air. (e) By heating chromic hydroxide in an inert gas, finally to a high temperature, the dark green exide first produced, glows and turns black and

rend 1880, 90, 1359) (2) Crystalline.-Preparation: (a) By fusion of the amorphous oxide in an electric furnace (Moissan, 181d. 1892, 115, 1035). (b) By heating potassium dichromato with aodium chloride (Ditte, 151d. 1902, 134, 336), with tin (Prod'homme, Bull Soc, Mulhouse, 1889, 59, 599). It has been observed as a kind of

becomes ansolublo in acids (Moissan, Compt.

aublimate in pottery furnaces. Wöhler prepared this oxide in fine, small rhombohedral crystals by passing the vapour of chromyl dichlorido through a tube heated to redness. The crystals are isomorphous with coundum and of equal hardness; their ap gr. 1852l.

A fine ahede of amorphous chromic oxide is produced by beating mereurous chromste Hg,CTO, in a covered crucible; mereury and nxygen escape, and the oxide remoins as o green powder. For the preparation of this aubstance on the large ocale, a great number of methods are recommended; nne early method is as follows: -boil a colution of potassium dichromato with half ats weight of flowers of sulphur so long as the green hydroxide is precipitated. The addition of a little potash colution, by forming potassium sulphide, accelerates the decom positing. The precipitate is filtered and washed. The sulphur retained in the precipitete may be removed by heating (Lassaigne, Ann, Chim. Phys. 1861, [ut], 14, 299); cf. Siegle & Co., B.P. 461799 and Addn. B P. 461800; Wöhler (Pogg. Ann. 1827, 10, 46) heats a mixture of potassium dichromate with its own weight of ammonium chingde and a small quantity of audium carbonate, and purifies the residue by washing According to Barian (Rev. Scient, 1846, 20.

fine porcelain, is produced by igniting in a crucible a mixture of 4 parts of potassium chromate and I part of starch The mass is washed free from potassium carbonate and re-

Chromium green is manufactured together with alkaline earth and alkali formates by licating a suspension or a solution of an alkalineearth or alkalı eliromate at 150°-350° with carbon monoxide under pressure (I G. l'arbenind

A G, BP, 305588).

Chromic sesquioxide is a green pigment of great permanence. It is not acted upon by chlorine or sulpliur gases or by an intense heat It melts at 1,990° (Kanolt, J. Washington Acad. Sci. 1913, 3, 315), at 2,140°±25° (Bunting, Bur. Stand J. Res 1930, 5, 325), and crystallises on cooling a chromium hydroxide (t. p. 108).

or Cr₄(OH)₁₀O,9H₂O (Jovitchitch, Helv. Chim Acta, 1920, 3, 46), absorbs earbon dioxide from the atmosphere until a saturation limit is reached The compound appears to be

$$CO_3[2Cr_2(OH)_6],8H_2O,$$
 $Cr_2(OH)_5\cdot O$
 $CO,9H_2O$
 $Cr_2(OH)_5\cdot O$

It can be dried at 100° without losing CO₂, which is, however, liberated by acids (Compt.

rend. 1914, 158, 872).

The oxido is largely used under the names chrome green (mixtures of chrome yellow and Prussian blue are also called chrome greens and must not be confused with the true chrome green) and ultramarine green for imparting a green colour to glass, porcelain, etc. It is used for producing the chrome red glaze in ceramic ware, as a pigment in oil and water colours and in printing, and as a mordant in calicoprinting and dyeing Certain hydrated oxides are also used under various names, their colours are, generally speaking, brighter than that of the anhydrous oxide, but they usually contain small quantities of other substances besides the oxide and water. (For hydrated chromic oxide i. R Fricke and G. F. Huttig, "Hydroxide and,

Oxylydrate," Academische Veilag Leipzig, 1937).
Gulgnet's Green; Pannetier's Green,
Emerald Green, Verdian, 2Cr₂O₃ 3H₂O.— This pigment appears to be identical with that formerly manufactured in secret by Pennettier According to Guignet's method, 3 parts of borie acid and I part of potassium dichromate are he sted to dull reduces in a reverberatory furnace The mass swells up, evolves oxygen, and becomes of a fine green colour, it contains the borates of potassium and chrominiu, or a double borate of those two metals. It is boiled with water, whereby the borate of chromium is decomposed into boric acid and hydrated chromium sesquioxide, potassium borate also remaining in colution. The precipitate is well washed, dried, and finely ground. It usually contains boric acid even after thorough washing, but not if prepand with ammonium dichromate The washincs and mother liquors are evaporated for the recovery of the boric neid

According to Chem.-Ztg. 1885, 9, the process then used on the large scale was as follows: The borne acid was first purified by solution in hot water, treating with animal charcoal and recrystallising The crystals were dried in a centrifugal machine. The mother liquor could be employed three or four times for recrystallisation of fresh portions of acid, but became then too impure for further use. Eight parts of potassium diehromate and 3 parts of purified borie acid were ground into a stiff paste with water, and the mixture in charges of about 1 ewt. heated to dull redness for about 4 hours in a reverberatory furnace. The fused mass was thrown into water, and washed repeatedly by decantation; the pigment was ground while wet, again washed, filtered, and dried. The first two washings contained considerable quantities of potassium borate, which was recovered by evaporating in leaden pans, adding hydrochloric acid and allowing the boric acid formed to crystallise. The mother liquor was further evaporated and crystallised. The crystals were redissolved, the lead (from the pans) precipitated by hydrogen sulphide, and the filtrate recrystallised. In this manner, from 70 to 75% of the borne acid was recovered.

Guignet's green is the most permanent green pigment known; it is not acted upon by light or concentrated boiling alkalis; it is not affected by acids in the cold, but hot hydrochloric acid slowly dissolves it. When heated to 200° it blackens and becomes anhydrous. It is a fino green pigment largely used for the same purpose as the ordinary oxide but is more brilliant, and may replace the dangerous arsenical greens. It possesses good covering power, and can be mixed with other pigments without alteration When mixed with lead chromate, it is employed for tho production of a pale green colour in landscape

painting.

Other chrome greens are obtained by the use of sodium phosphate. These always contain some phosphoric acid. They are, however, not as brilliant as the oxide chrome greens. Various methods of preparation are employed. Arnaudon dissolves ammonium phosphato with slight excess of potassium diehromate in a little boiling water, evaporates until the mass solidifies on cooling, heats at 80°, and later at 200°, washes with hot water, dries, and powders Another method consists in boiling a solution of 10 lb. potassium dichromate with 18 lb sodium phosphate, adding 10 lb. sodium thiosulphate and a little hydrochloric acid, and boiling until precipitation is complete The precipitate is washed, ground, and dried.

Blanc and Chaudron (Compt. rend. 1926, 182, 356) state that the blue form of Cr_2O_3 changes

to green at 500° m air, and then resembles Guignet's green (cf B, 1926, 166).

Chromous Salts.—Trube and Goodson (Ber. 1916, 49, 1679) have prepared chromous salts by the electrolytic reduction of the corresponding chromic salts at a cathode of pure The most favourable conditions for the reduction of vic let chromic salts are in moderately acid concentrated solutions with a current density of 25 amps per sq decimetre. The more common green salts can be used in more

with doubla or triple linkings in the presence of water. Acetylene is reduced to ethylene, but not to ethane; maleic and fumane acids are readily reduced to encuric acid. Nitrous oxide in the presence of alkalı hydroxide is reduced to nitrogen. Nitro acid and hydroxylamine are reduced quantitatively to ammonia (Traube and Passarge, Ber. 1916, 49, 1692). Chromous salts form stable double compounds with bydrazine which may be precipitated from chromous acctate dissolved in air-free water covered with light petroleum Chromous diby drazine chloride, CrCl, 2N,H, is a hlac coloured powder sparingly soluble in wate.

M. Chatelet (Compt rend 1934, 199, 290)
M. Chatelet (Compt rend 1934, 199, 290)
prepared anhydrous CrCl, by reduction of
CrCl, with H, at about 700° It dissolves in
C,H₃N, yielding a green solution and green
crystals of the compound CrCl, 2C,H,N. This
compound abows traces of oxidation effer some hours in the air.; I mol of CrCl₂ absorbs 548 mols of dry NH₂ at atmospheric pressure and at room temperature higher absorption found by Ephraim and Zapata is attributed to the lower reduction tempera ture. The composition varies with ΝH. pressure and at zero pressure corresponds with CrCl,3NH, confirming the existence of this compound, but no evidence could be obtained compound, but he evidence count to obtained of the existence of CrCl₂5NH₂ by studying the pressure composition curre. When, however, the CrCl₂ is kept asturated with NH₂ at 22° and 76 cm, the colour changes from pale green to violet rose, and in 20 hours the product corresponds with the composition CrCl, 5NH, and its composition does not vary over changes of NH, pressure from 11 to 83 cm.

Chromic Hydroxide is precipitated from a solution by caustic alkali or ammonia, but can only be obtained pure by use of the latter reagent owing to adsorption of the fixed alkali by the precipitate. The water retained varies

with the method of drying

A V. Rakovski and T. Poljanski (Trans. Inst. A V. Rakovski and T. Poljanski (Trans. Inst. Pure Chem. Reag. U.S.S.R. 1931, No. 12, 11-16) find that Cr. hydroxide free from Cf may be obtained by reducing Na₂Cr₂O₂ with CH₂O in presence of HCl, precipitating with NH, and washing with dilute NH2 and then with H2O.
Preparations made from the sulphate cannot be freed completely from SO, linhn (Ber. 1932, 65, [B], 61) recommends 15 g NaNO, 15 g. NaN₃ in H₂O (250 c.c.) as a reagent for obtaining a pure, dense, easily filtered form. A colloidal solution of hydrated chrome oxida

was prepared by Graham (Phd. Trans 1861, 151. 183) by dualysis of a solution of freshly precipitated chromio hydrovide in chromie chloride solution in water After 30 days' dialysis the deep green solution showed 43 parts HCI to 95 7 parts Cr2O2, after 38 days 15 acid and

concentrated solutions, but require a greater density of current.

Anhydrous chromous salts are white and yield blue solutions which are gradually oxidated and yield blue solutions which are gradually oxidated and the solution of CrX, in the etner R,O. They are on that SCPO+H,O-Cr,O,+H, They are on this account able to reduce negation compounds of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when NH, is added to a representation of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when NH, is added to a representation of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when NH, is added to a representation of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when NH, is added to a representation with the highest processing the compounds of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when NH, is added to a representation with the highest processing the compounds of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when NH, is added to a representation of hydrogen are precipitated when the time of the processing that the compounds of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when the processing the compounds of the type CrX, OR,SNH, Where X is a hadogen and R an alkyl radical are precipitated when the type CrX, or the processing the proc

J. Laestecks (Rocz. Chem. 1930, 10, 736) finds that stable chromic oxide sols are obtained by adding washed chromic bydrouds prepared by the action of ammonia on chromic netrate aclution to a solution of chromic chloride

such sols do not need to be dialysed. Chromic Chioride, CrCI, or Cr,CI, anhydrous chlorida is obtained by heating mixture of carbon and chromium sesonoxide in a current of chlorine. It can be conveniently prepared by Bourion's method (Compt. rend. 1909, 148, 170) by passing the vapour of sulphur chloride, bp. 138', over the oxide gradually raised to a red heat. It forms pale-violet scales of sp gr. 2 782 (Bilts and Birk, Z. snorg. Chem. 1924, 134, 123); 2 916 (Crespi, Anal, Fis. Quim. 1928, 28, 182); it is almost insoluble in water, but dissolves readily, being transformed to the green variety (v anfra), if only a minute trace of chromous chlonde (1 in 20,000) is present,

Chromic chloride may be prepared in solution by dissolving the hydroxide in hydrochlone and The hydrates of chromic chloride include three isomene hexahydrates : a green salt,

[CrCl.(H.O),]Cl.2H.O.

a violet salt, [Cr(H₂O)₆]Cl₂ (Recours, Compt. rend 1886, 102, 548; Ann Chim. 1887, 10, 34). rend 1950, 100, 001 Ann Culm. 1951, 10, 50, and a pale green aslt of intermediate constitution, [CrCi(H₁O₂)Cl₂H₁O [Bjerrum, Ber 1906, 39, 1899; Z. physikal (hem. 1907, 59 376, 896). The green hexahydrato is parly dehydrated in some over sulphuric and see yields the tetrahydrate, a pale green pooler, chromic dichlorotetraquochloride,

[CrCl_(H_O),]Cl,

The green decahydrate.

[CrCl_(H,O),]Cl,6H,O,

crystallises from a strongly cooled solution (66%) of the green hexahydrate (Werner and Gubser, Ber. 1906, 39, 1823). The latter salt kept for 4 months at 1 mm. over phosphorus CrCl, H.O is formed.

The formulæ of Werner and Gubser (lc)

explain the behaviour of the green hers hydrate (supra); in a freshly prepared solution at 0°, only one third of the chlorine rescu with silver nitrate. At ordinary temperature the AgCl precipitat may correspond to all values between 14 and 3Cl. The solution of the value between 14 and 3Cl. The solution of the violet beachydrate yields 3AgCl. Hantzed and Torke (Z. anorg. Chem. 1932, 299, 60) prepared a bright bluish-green hydrate I from or parts Cr.O., after 38 days 15 and and [Cr(H,O),Cl., and a dark green hydrat 18 5 Cr.O. [Cr(H,O),Cl., and a dark green hydrat liften [Cr(H,O),Cl.,Cl.,2H,O. by the addition of NH,OH and NH,Cl. (1) is readily soluble in acids, (II) less soluble. The formulæ suggested reacts vigorously with pyrophoric Cr at 300°, are (I) Cr(OH)3,3H2O and with partial formation (67%) of Cr13.

(II) [Cr(H₂O)(OH)₃],2H₂O.

On passing hydrogen ebloride through a green solution of ebromie chloride a helietrope coloured double chloride, CrCl, HCl, xH,O (x approx. 6) is precipitated. The substance is insoluble in methyl and ethyl alcohels, and in water gives a pale red solution which immediately turns green (Partington and Tweedy, J.C.S. 1927, 2899).

Chromyl Dichloride, CrO₂Cl₂.—Moissan (Compt. rend. 1884, 98, 1581) prepared chromyl chleride by treating dry chromic anhydride with hydrogen chloride. It forms at the ordinary temperature but the action is accelerated by heating. It may also be made hy distilling petassium bichromato (3 parts) and common salt (3 parts) with sulphuric acid (9 parts) in an atmosphere of carbon dioxide. The corresponding bromide and iodido cannot be made hy this method, but the fluoride can, the formula for the latter being CrO₂F₂ (Gazz. chim. ital. 1886, 16, 218). H. D. Law and F. M. Perkin (J.C.S. 1907, 91, 191) found that the hest way of preparing this compound is to dissolve chromie anhydride (50 g.) in strong hydrochloric acid (170 c.c.) to which is added 100 c.c. of strong sulphuric acid, 20 c.c. at a time. Chromyl chloride sinks as a red liquid and may be separated in a tap funnel. B.p. at 760 mm. is 116.7°, m.p. -96.5° ±0.5°. It is a deep red liquid resembling bromine. Two specimens had d_{4}^{-47} , 2.0528-2.0515; d_{4}^{0} 1.9591-1.9582; d_{4}^{25} 1.9124-1.9113 (Moles and Gomez, Z. physikal. Chem. 1912, 80, 513). Its vapour is not dissociated at 181°C. When heated in a elesed tube to 180°-190° trichromyl chloride, (CrO2)3Cl2, is formed. This is a black, noncrystalline powder deliquescing in air. Chromyl dichlorido reacts with phosphorus trichloride in earlion tetrachloride solution to form Crocl, Pocl, and Pcls.

Chromous Bromlde, CrBr2.—A white salt giving a blue solution, prepared by Moissan by passing bromine vapour with nitrogen over heated chromium.

Chromic Bromlde, CrBr3, is formed like the dibromide, hy passing excess of Br over heated Cr. It is a dark green powder and

ferms two isomerie hexahydrates, green and violet, corresponding to the similarly coloured

hexalivdrates of CrCl3.

Chromous lodlde.—Hein and Wintner-llolder (Z. anorg. Chem. 1931, 202, 81-89) pre-pared pure Cri₂, m.p. 790°-795°, by heating electrolytic Cr with an excess of 1 in N₂ or vacuum at 1,150°-1,200°, and removing the excess of 1 from the product by heating at 200°. Two forms viz growich white and death howe Two forms, viz., greyish-white and dark hrown, exist. Crl, has an appreciable decomp. pressure at 400°-700°, but in presence of a trace of I vapour this is strongly depressed, and increases only very slowly with rise of temperature. Iodine reacts, but not very readily with Crl. at 300°, a product containing more air-free water covered with a layer of light than 91% Crl. could not be obtained. It petroleum. The acetate is dissolved by addition

Chromic Fluoride, CrF2,4H2O.—This product, prepared by dissolving chromic hydrate in hydrofluoric acid, is used in the printing and dyeing of woollen goods. (For preparation, v. 1937, B, 1199). It is a fine, crystalline, green powder, very soluble in water, but it bas a corroding action on glass, and is therefore best kept in lead or wooden vessels. It may be used as a mordant with both vegetable and animal fabrics, and is often employed in the place of acctate or nitrate of cbromium for printing, and, in general, gives finer results. Alizarin dyes, cœrulcin, gallein, etc., and logwood extracts, are fixed as perfectly as with acctate of ebromium. One advantage is that it is marketed as a solid. Its effect on wool in after-chroming baths bas been criticised (Knecht, Lawson, and Loewen-thal, "Dycing," II, 632, London, 1925). Chromium silicofluoride, prepared by dissolving tho oxide in hydrofluosilicie acid, bas also been used in dyeing and printing in place of the fluoride, and is stated to give better results.

Chromium Nitride .- Valensi (Compt. rend. 1928, 187, 293) prepared chromium nitride as a brown crystalline powder by heating chromium in pure nitrogen. It gives dissociation isotherms between 810° and 1,200° analogous to those of palladium and hydrogen, which indicate the formation of the compound CrN. The temperature, 1,015°, at which the dissociation pressure is 760 mm. agrees with the calculated

value.

Chromium compounds with nitrogen prepared by passing ammonia over chromium heated to 800° show the existence of two phases. The first phase is homogeneous for a considerable rango of concentration which probably includes 33 atomic per cent. nitrogen corresponding to tho formula Cr₂N. The chromium atoms are arranged in a hexagonal lattice with the densest spherical packing, the nitrogen atoms probably being distributed at random in the spaces of the lattice. The second phase corresponds to tho formula CrN with NaCl structure. X-ray examination of a nitrogenous ferrochromo (60.3% Cr, 2.4% N, 0.7% Mn, 2% Sl, 0.2% C) showed hexagonal CrN (Blix, Z. physikal. Chem. 1929, B, 3, 229).

In the system Cr-N the hexagonal \(\beta\)-phase bas a "superstructure," the cell volume of which is three times as large as that of the hexagonal closc-packed lattice. The upper and lower limits of the region of homogeneity of the B-phase are 11.9% N (Cr₂N) and 9.30% N respectively (S. Eriksson, A, 1935, 1455).

Chromium Azlde, Cr(N₃)₃ (Mandala and Comella, Gazzetta, 1922, 52, (i), 112) was prepared by prolonged treatment of an absolute alcoholic solution of crystalline chromium nitrate with sodium sulphate, and evaporation of the filtered liquid with the calculated proportion of sodium azide in a vacuum. It forms a highly hygroscopic dark green amorphous mass.

Chromous Hydrazine Compounds (Traube and Passarge, Ber. 1913, 46 1505) can be pre-pared by suspension of chromous acetate in of the exactly necessary quantity of dilute solution has an alkaline reaction. Like an acid, and then a solution of hydrazinn hydrate or sulphate edded. The salt precipitates readily and is washed with water, alcohol, and ether. and is washed with water, alcohol, and ether. Chromous dihydrazine chloride, CrCi₂,2N₂H₂, and the hromide, CrBr₂,2N₂H₃, are hias-coloured compounds, the lodde, CrI₂,2N₂H₃ is grevish-blue

Chromium Nitrate.-Jovitschitsch (Mon atsh. 1912, 33, 9-18), hy dissolving atrongly heated chromic oxide in hot concentrated HNO. (d. I-4), obtained a solution from which this hydrate, Cr₂(NO₂)₄,15H₂O, erystallises on ecolog in the form of dark hrown prisms (a.h.c.=1.4210:1:11158, β=93° 10') cootact with dry air the ervstals lose 6H.O with the formation of a grey coloured hydrate, Cr. (NO.) 9H.O Partington and Tweedy by

dissolving chromic hydrate obtained the salt Cr. (NO₃)₆, 25H₂O (J C S. 1926, 1142) Chromium Silicides,—Chromium silicides

may be prepared by heating the two elements mixtures of chromic oxide and silicon carbide in the electric furnace, compounds represented by SiCr₂, SiCr₂, Si₂Cr₃, and Si₂Cr, have been prepared They are crystallino and very hard—SiCr, is harder than corundum—and are not attacked by ordinary acids, though attacked readily by hydrofluoric acid and by equa regia (Lebeau and Figueras, Compt rend. 1903, 138, 1329, Matignon and Trannoy, ibid. 1905, 141, 190, Frilley, Rev. métal 1911, 8, 476, Bareduc Muller, ibid 1909, 6, 157; 476 , Barec 1910, 7, 698)

B Boren (Arkiv Kemi, Min Geol 1934, 11A, No. 10, 23 pp) inferred the formation of the following compounds in the system Cr-Si. Cubic crystals of Cr-Si with a 4 555A and 8 atoms in the unit cell, a phase of uncertain composition and atructure stable only at about 1,000°; CrSi, cuhic, a 4 620Å, apece group T4, S ctoms in the unit cell, atructure analogous to that of FeSi; CrSi, hersgonal, a 4 422,

s 6 351A , space group D.

Chromium Borides, - Moissan prepared hard emorphous CrB from the two elements in the electric furcace (Ann. Chim. Phys. 1896, [vn], 8, 565). Wedekind and Fetzer employed a mixtore of chromic oxide, aluminium powder, and powdered boron, and obtained crystalling and powdered boron, and obtained crystalling UrB (Ber. 1907, 40, 297) Andricux prepared Cr₂B₂ by electrolysis of a fused mixture of B₂O₃, Cr₂O₅, MgO, and MgF₄ (Ann chim 1929, [x], 12, 489).

Potassium Chromate, K, CrO, 18 prepared, as already described, from chromn ore. It may be produced by adding potassium hydroxide to a solution of the inchromate. It crystallises in anhydrous, yellow, rhombie prisme isomorphous with those of potassium aulphate, sp gr. 271 (Kopp). 100 parts of water dissolved of potassium chromate

0° 30° ഗ്ര 105.82 57-11 65 13 74 60 **88 8**

105 8° being the hoiling-point of the satureted solution (hoppel and Blumenthal, Z. anong and retains the pigment, while the portions from 1907, 53, 262)

The solution has a fine texted by the darker parts of the negative are

chromates the salt is poisonous (r. Chromes supra).

When heated, it reddeos and fuses without decomposition; heated with reducing agents

cuch as sulphur, it is reduced to chromium sesquioxide. The addition of an acid, even carbonie acid, determines the decomposition into potassium dichromate. It forms double salts with the chromates of the alkahoe carths and

Potassium chromate is not largely used, the dichromate, on account of its higher percentage

of chromic acid, belog geoers lly preferred.
Potassium Dichromate, K₂Cr₂O₂, mp
389 4°±0 5°.—This important salt, generally known as "bichromate," or "hichrome," is prepared in large quantities by the methods already described. Considerably over 10,000 tons are annually produced in Great Britain

It erystallises in anhydrous, red, tabular prisms belonging to the triclinic system, of sper 270, and of intensely metallic, bitter taste At bright redness oxygen is evolved, and the normal chromate and chromic sesquioxide result Acid chromates, of composition K2Cr2O10 2nd K2Cr4O13, appear to exist in solution within certain limits of concentration.

100 parts of weter dissolve et 0°, 4 64 parts; at 30°, 18 13 parts; at 60°, 45 44 parts; and at 104 8°, 108 2 parts, 104 8° being the boiling point of the caturated solution (Koppel and Blumenthal, Z anorg Chem 1907, 53, 263)

The salt separates from the fused state as a compact mass of dark brownish-red tabular crystals with marked reduction in volume On further cooling the crystels change to e loos, orange-red powder, the transition being 236 5'z The unstable monoclinic form changes to stable tricking crystals et this temperature (Robinson, Stephenson, and Briscoe, JCS.

1925, 127, 547). Potassium dichromate is used in the prepara-tion of chrome pigments, of "discharge" for Turkey red, etc., for the production of a large variety of colours in colico printing and dyeing, and in the manufacture of safety matches In colution with sulphuric soid, it is used as a bleaching agent for tallow, palm oil, etc., in the oxidation of anthracene to all comm. in the manufacture of milian with and in many other organic oxidations also used in the two bath process of chrome tanning. The pelt is immersed first in the dichromate solution and then in a solution of

sodium thiosulphate acidified with HCi. When mixed with organic substances it is reduced on exposure to light; gelatioe in such circumstances is rendered insoluble. This is action is taken advantage of in the "carbon The geliure process of photographic printing. is mixed with a pigment of any colour, and the paper carrying this film is sensitised by floating on a solution of potassium dichromate. posure under a negative, the gelatine in those portions exposed to the light becomes insoluble yellow colour, e distinct yellow tinge is imparted almost unacted upon, and may be dissolted r by 1 part in 400,000 parts of water. This warm water. In this manner, photographs of

may be produced.

In the oil-process gelatine-coated paper may be sensitised with bichromate and exposed under a negative and, after developing the print in water at 25°, it is pigmented with a suitable oil-ink, since the hardened gelatine has the property of fixing greasy inks. In the corresponding gum-bichromate process the paper is coated with a solution of gum stable and bichromate. For the carbro process a bromide print is soaked in water and squeegeed into contact with the sensitive tissue of the carhon process. The silver grains of the print reduce the bichromated gelatine, rendering it insoluble, so that after separation of the tissue it may be developed in hot water. In the bromoil process a hromido print is bleached in a bichromate bath, whereby the gelatino of tho silver image becomes insoluble, fixed, washed, and the hardened gelatino pigmented brushing on an oil-ink as in the oil-process.

Sodium Chromates.—The normal chromate Na, CrO4,10H2O, and the diehromato

Na₂Cr₂O₇,2H₂O,

may be prepared by methods similar to those already described under Chromates (q.v.), but without the details which now follow. According to Nie. Walberg (Dingl. poly. J. 1886, 259, 188), 6 parts of powdered chromo ore (44% Cr₂O₃) were mixed with 3 parts of soda ash (92% Na₂CO₃) and 3 parts of chalk, and heated in charges of I ton in a reverberatory furnace for 8 hours. The mass is lixiviated to produce a solution of 45°B., boiled down to 52°B., and allowed to crystallise in leaden pans. crystals are first dried by a centrifugal machine and finally heated to 30° in a drying chamber, where they crumble to a yellow, anhydrous powder centaining about 96% of the normal chromate.

For the production of dichromate, these crystals are dissolved to a solution of 40°B. and treated with sufficient chamber acid to determine the conversion into the dichromate, the right point being found by potassium iodide and starch paper. Sufficient neutral sodium chromate is then added to bring the percentago of CrO_3 to about 72.5. The liquor is cooled in lead tanks to 1°, and the sodium sulphate crystal-The liquor is drained off, filtered if necessary, and evaporated to dryness in an iron The residue is pot, with constant stirring. powdered while still hot. If the amount of CrO₃ be allowed to exceed 72.5%, the product is damp and cannot be stored in wooden casks. An analysis of this substance showed CrO₃ 72-3, Na O 26-20, SO 1-40.

Modifications in the method of manufacture by Ulm (Chem.-Ztg. 1914, 38, 670; J.S.C.I. 1914, 33, 917), consist in the use of caustic alkali in the place of alkali earbonate, and in the employment of only half the sodium carbonate required to decompose the ore, the remainder being replaced by sodium sulplinte, formed in the later stages of manufacture.

Normal Sodium Chromate forms large, efflorescent erystals of the Na₂CrO₄,10H₂O, isomorphous with Glauber's organic analysis.

any desired colour, and of great permanence, | salt. Unliko potassium chromate, it is less solublo than the dichromato; it is, however, more solublo than potassium chromate, tho saturated solution at 18° contains 40·10% Na₂CrO₄ (Mylius and Funk, Ber. 1897, 30, 1718).

Other hydrates with 6 and 4 mols. of H.O aro known. Their transition temperatures have been suggested as convenient fixed points in thermometry (Richards and Kelley, J. Amer.

Chem. Soc. 1911, 33, 847).

Sodium Dichromate, Na, Cr, O, 2H, O. crystallises in thin, six-sided prisms of a fine hyacinth-red colour, and is very solublo in water, 100 parts of water at 30° dissolving 197.6 parts Na₂Cr₂O₇ (Schreinemakers, Z. physikal. Chem. 1906, 55, 91). It is hence more than 10 times as soluble as potassium dichromate (18.12 parts), which it frequently replaces. (For a difference in properties, see Rohinson et al., l.c.)

Ammonium Chromate and Dichromate are prepared by mixing in proper proportions solutions of ammonia and of chromic acid. The dichromate, on heating, yields water and a mixture of equal volumes of nitrogen and nitrous oxide and leaves a residue of chromic oxide mixed with a bigher oxide of chromium

(Harbord and King, l.c.).

Ammonium chromate may be also manufactured by treatment of sodium chromate solution with ammonia and saturating with carbon dioxide, whereby sodium bicarbonate is

precipitated.

Calcium Chromate, CaCrO4,2H2O, is largely produced in the manufacture of chromates from chrome iron ore. The dihydrate exists in monoclinic α and rhombie β forms. The anhydrous salt formed at 200° is less

soluble at 100° (0.42 g.) than at 0° (4.31 g. in 100 g. solution) (Mylius, Ber. 1900, 33, 3689).

Barium Chromate, BaCrO₄, is a canary-yellow powder, known as Yellow Ultramarine or Lemon Yellow. It is produced by the addition of a chromate to a solution of a salt of harium.

Barium chromate is insoluble in water and acetic acid, solublo in hydrochloric and nitric acids. On treatment with a boiling solution of chromic anhydride, it dissolves, and crystallises on cooling as a yellowish-red powder consisting of the dichromate BaCr2O7,2H2O.

Both barium and calcium chromates have been employed as pigments, but are now little used on account of their lack of hrightness and

covering power.

Lead Chromates.—The neutral lead chromate PbCrO₄ is best produced by the action of a solution of lead acetate on a solution of a chromate. Thus produced, it is of a fine lemonyellow colour, insoluble in water and dilute acids. At 25° the solubility of PbCrO4 in water is 5·8×10-5 g./l. (Ishibashi and J. Chem. Soc. Japan, 1936, 57, 1028). When heated at 250° it becomes reddish brown, at a higher temperature it fuses, and finally evolves oxygen with the formation of chromic sesquioxide and a basic lead chromate. On account of this evolution of oxygen the subcomposition stance is frequently used for combustions in

caustic alkalı, but is not dissolved by hut lime water. For this reason, in immersing calico K, SO, and PhCrO, and that the existence of coloured with chrome yellow in lime water, double salts is inhikely. The strong adsorption for the production of an orange colour, care rendered the determination of the mass-action abould be taken that the solution is bot.

When heated with the calculated amounts of caustic alkalı part of the cbromic acid may be removed, with the production of basic lead chromates of colour varying from urange to vermilion.

The normal chromate is largely used as a pigment, and for calico-printing, and is the essential constituent of various chreme mg ments as chrome vellow, Paris vellow, and Leipsic vellow. The nama Cologne vellow was originally given to chroma yellow containing a mixture of lead chromate and lead sulphate, but the latter compound is also present in most of the bright Jellows known commercially as "genuina"

chroma vellowa Chrome Yellows .- The finest chrome yellow ia produced by precipitation; the lead acetate used should not be basic or the product will be of an orange colour: the solution should be dilute, and the lead salt should be in excess, the presence of excess of chromate is liable to "turn" the yellow, se to produce on orange tingo. The substances used should be technically pure, According to BP 416744, 1933, one method of manufacture from chromic acid consists in adding this acid to a suspension of lead oxide or carbonate in hydrofluoric, fluoberic, or fluosilicio acid, at a rate which is not greater than the velocity of reaction of the latter acids Kohn found that the brightest chrome yellow with the lead compound Pure chrome yellow, PbCrO, is too expensive for most purposes, is hable to darken in colour, and does not exhibit the required range of tones. A series of "genuine" chrome yellows is manufactured by mixing the precipitating chromate solution with an appropriate quantity of sulphurse acid, Glauber's salt, or alum, the latter salt for American chrome-yellow. The lead chromate is thus precipitated together with any required amount of lead sulphate as mixed crystals of PhCrO, and PhSO, the colour being paler as the proportion of lead sulphate is mereased. range of "reduced chroma yellows" is made by the admixture of barytes, gypsum, keelin, or whiting, which are ground with water and the suspension added to the precipitating tank prior to the reaction. As the customer requires chroma jellows, eg. primrosa chrome, lemun chrome, and middle chrome, which for reasons uf shade or price contain lead sulphate or fillers. these additions are not in general to be regarded as adulterants. Lemon Chroma A and Orange Chroma B may contain from 25 to 90% barytes. The application to these chromes of the Hume Office test for toxicity has been studied (Samuels, J. Oil Col. Chem. Assoc. 1938, 21, 177). The British Standard specification for lead chromes for paints is 282/1927.

Milbaner and Kohn (Z. physikal Chem. 1916,

of left to right. From determinations of the green pigments, and the fastoess to light of such

Lead chromate is soluble in cold lime water or stability it is established that the solid phase PbSO, cannot exist in the presence of K,CrO, curve difficult. The general results indicate that the manufacture of chrome yellows can ba carried outfromleadsulphate and that various shades can be obtained by varying conditions, temperature, concentration, and time. It follows also that an excess uf lead acetate should be used in the manufacture, since this will secure the absence of K2CrO4, which would otherwise rapidly con-

ticular abade into chromate, Freshly preceptated PhCrO, darkens in colour probably because of the change from an unstable rhombie into a stable monochine form. When lead sulphate is precipitated amultaneously with lead chromata the two salts form mixed monochnic crystals which do not darken during settling. The best conditions for precipitation were ascertained by Wagner and Keddel (Farben Zig. 1926, 31, 1567) Tha monoclime mixed crystals of PbCrO₄ with PbSO₄, BaSO₄, CaSO₄, or SrSO₄ are said to be more resistant to light than the rhombto forms and the rhombic mixed crystals first precipitated may be converted into monoclinio beating their aqueous suspensions (BP. 403762) Precipitation from dilute solutions with vigorous stirring produces finer particles and brighter shades, while redder tones are obtained in hot solutions. Milhauer and was obtained by adding a solution of lead mirate or acctate to a fillute solution of potassium chromata and potassium sulphate (J.S.C.I. 1923, 1954; Chem. Ztg. 1922, 48, 1145). They showed that the precipitate consisted of rhombic mixed crystals. These mixed crystals were shown to be solid solutions by X-ray examination (Lederle, Rev. Prod. Chim. 1937, 40, No. 9, 263). The monoclinic form is asid to be superior to the rhombio in resisting light (Wagner, Paint and Varnish Prod. Man. 1934, 10, No. 5, 10).

The method usually adopted for production of chrome yellow is the following: The two precipitating solutions are prepared asparately in wooden tanks and heated by steam pipes. Tha solutions are either filtered or allowed to settle in the tanks and drawn off from above. They are run together into the precipitating tank, which contains about 200 gallons. The chrome sellow is allowed to deposit and the clear liquid drawn uff, more water is added, the whola thoroughly stirred, and the process is repeated three or four times. The pigment is then pumped into a filter press and dried in drying rooms at 30°-50°.

Muxtures of chrome yellows and Prussian blua form the Brunswick greens, also known as chrome greene, although they are to be distinguished from the more expensive and 91, 419) show that the reaction
PbSO4+K4CrO4⇒PbCrO4+K4SO4
proceeds practically completely in the direction
Blue has been mixed with chroma yellow for 187). By the regulated action of bot alkalis on chrome yellow a series of basic lead chromates is produced ranging in colour from orange chrome to chrome red.

Orange Chrome.-A washed chrome yellow precipitato consisting of PbCrO4 and PbSO4 is treated at 80°-90° while stirring with the calculated quantity of caustic soda solution until the reaction is completed. The product is washed, filter-pressed, and dried as usual (Zerr and Rübencamp, "Farbenfabrikation," 4th ed., Berlin, 1930); or the freshly precipitated lead chromate+lead sulphate is boiled with milk of lime-with the object of producing a basic chromate containing gypsum as a filler. In another process basic lead chloride is first prepared by the action of hot sodium obloride solution on litbarge, dichromate solution is then added which forms monochromato with the caustic soda now present and this reacts with the basic lead salt. Orange chromes sometimes contain a colour lake precipitated on the pigment to brighten the shade. Holley ("Analysis of Paint and Varnish Products," p. 214, London and New York, 1912) found 4.87% of organic and New John, 122 told 48.7% of diginal colour in a pigment containing PbCrO₄ 40.56%, PbO 47.24%, PbSO₄ 5.49%, which may be compared with a light yellow chrome containing PbCrO₄ 68.65%, PbSO₄ 31.21%. Chrome Red is conveniently manufactured by

boiling white lead with a solution prepared from dichromate and caustic soda, the washed product which has a dull appearance acquires a brilliant red colour after treatment with diluto sulphurio acid, the pigment is then washed and dried as usual. The pigment is not so generally useful as the other lead chromes, for when finely ground the shade changes from deep red towards orange yellow. It may replace einnabar in distemper painting, but settles out too readily if made into an oil paint. The synonyms of this pigment include the names Derby red, Persian red, American vermilion, Victoria red, Chinese red.

Wagner and Schirmer (Z. anorg. Chem. 1935, 222, 245) studied the products formed by adding KOH to a lead salt and potassium chromate. With increasing KOH the orange red PbO·PbCrO4 changes to a pure red substance of the same composition and crystalline form (tetragonal) but of different particle size. The final product was orange red, not uniform, and appeared to contain yellow PbO.

Holley (I.c. 220) found in a light American ver-

milion, PbCrO₄ 50·16%, PbO 41·20%, PbSO₄ 6.15%; in a deep American vermilion PbCrO₄ 53.60%, PbO 40.88%, PbSO₄ 4.97%. It may be noted that PbO.PbCrO₄ contains PbO 40.85%, PbCrO₄ 59.15%. For rust prevention, a priming coat of a basic lead chromato paint (which may have an addition of zine chromate) has given excellent results on properly prepared structural iron and steel work. Chrome yellow paints darken when exposed to the air of towns and are also liable to be darkened by strong sunlight owing to the reducing action of the oil medium on the chromato (Palmer, J. Oil Col. Chem. Assoc. 1925, 8, 90). Reddy Britton and Evans (J.S.C.I. 1936, 55, 337T),

mixtures has been discussed by Amies and by pared the different types of lead chromes and Samuels (J. Oil Col. Chem. Assoc. 1938, 21, zinc chromes as regards grinding, hiding power, paint media, anti-corrosivo action, etc. The importance of the yellow and orange chrome pigments is shown by the output in the U.S.A. which in 1935 amounted to 39 million lb.

Scarlet Chromes is the descriptive name given to a series of pigments consisting of mixed crystals of chromate, sulphate, and molybdate of lead (J. Oil Col. Chem. Assoc. 1938, 174).

The lead chromates all possess good covering power and brightness of colour; they are durable, but blacken under the action of sul-phuretted bydrogen. They mix well with other pigments, but should not be used with ultra-

marine, lithopone or other sulphide pigments.

Blsmuth Chromates.—The addition of potassium dichromate to a neutral solution of bismuth nitrate gives a yellow flocculent pre-Bi₂O₃,2CrO₃, cipitate, soluble (Löwe). On heating, it becomes partly decomposed and turns greenish. K₂CrO₄ and solution of bismuth nitrate, the precipitate formed consists of 3Bi₂O₃,2CrO₃ (Löwe, 1856), and is quite insoluble in water. Bismuth chromate is used as a pigment (cf. Cox, Z. anorg. Chem. 1906, 50, 226).

Zinc Chromate, Zinc Yellow, or Buttercup Yellow, ZnCrO₄, is a beautiful yellow pigment, produced by the addition of hot neutral solution of zine sulphate to potassium chromate, or by adding a cream of zine oxide in water to a boiling solution of potassium dichromate. It is slightly soluble in water and readily so in nearly all acids, in ammonia, and

in caustie soda.

Zine chromate can be obtained in various tints and has a good colour. Though not equal to chrome yellow in covering power or body, it has the advantage of not blackening by contact with sulphides. It is used for mixing with Prussian blue to form the zinc greens, which are faster to light than the corresponding Brunswick greens. The British Standard specification for zine chromes for paints is 389/1929.

Samples of commercial zine yellow analysed by Fox, Ellis, and Hirst (J. Oil Col. Chem. Assoc. 1928, 11, 195; cf. ibid. 198) showed an average content of ZnO, 42.5%; CrO₃, 38.4%; theory for ZnCrO₄, ZnO 44.9%, CrO₃ 55.1%. The samples contained alkali, up to 11.1% K₂O, in another sample 7.0% Na₂O; and combined water up to 8.2%. Zing chromato purplessed from chemical supply Zine chromato purchased from chemical supply firms had a similar composition and in one case contained ammonia equivalent to 10.1% (NH₄)₂O. The potassium zine chrome above mentioned gave 15.1% water-soluble (arbitrary method in B.S.S. 254/1926), and the sedium zinc chrome 42.6%. In the discussion (l.c. 200), the change of colour in zinc chromate on keeping as a standard in a scaled bottle was mentioned, and also the loss on washing the pigment which could reach 50 to 60%. Brizzolaria et al. (Ind Eng. Chem. 1937, 29, 656) represent the composition by the formula

K₂O,4ZnO,4CrO₃,3H₂O.

(Official Digest, 1934, No. 130, 328) has com- reporting on seven years' exposure of protec-

tive painting on steel, found zinc chromate 1094, with bibliography, pp 1094-1098, Paris, priming paint very good, although not quite 1933. Duval considered that the 2,000 complex equal to red lead.

Briggs (J.CS, 1929, 242) confirmed the existence of normal zinc chromate and prepared the following crystalline chromates:

9ZrO, 5CrO, 12H,O, 3AI,O, 2CrO, 6H,O, 2Fe,O, 4CrO, H,O, 3SbO, CrO,

Perchromic Acid,-On adding hydrogen peroxide to an aggrous solution of chromie acid, or to an acucous solution of a chromate acidified 1924. with sulphuric acid, a deep blue coloration is produced, which on shaking the solution with

ether passes into that hould The reaction is a delicate test for CrO or H₁O₃. The blue perchromic acid is extremely unstable, evolves oxygen and is gradually Iransformed into chromie acid Alcoholic alkalies react with the ethereal solution forming salts of hise perchromic and, eg K₂Cr₂O₃₊, 2H₂O.Tr.Cr.O₁, (Schwarz et al. Ber. 1933, 66, [B], 310) The pyridine compound, however, 68. [B], 310) The pyridine compound, nowever, is (C₂H₃H)CrO₂ (bid 65, 871) Red perchromates, M₂Cr₂O₁₄, are formed by the interaction of alkaline chromate solutions and bydrogen peroxido below 0°, and the free acid is similarly formed from freshly precipitated chromio hydrale When scidified the solutions evolve oxygen and blue perchromic acid is formed. Bichwarz and Elstner (Ber 1936, 69, [B], 575) found the etherate CrO, Me,O to be (15), not hold the three to 0.3, was 0 to the product when CrO, reseted with roncentested H₂O₂ in Me₂O at low temperatures, this explodes already at -30°. The third series of perchromates, the brown salts, coordination compounds of the oxide CrO., include perchromateammine, CrO4,3NH, formed when H₂O₃ sets on an ammoniacal solution of ammonium chromate. The blue salts were formerly ronsidered to be MCrO, H,O, (Wiede, 1897-1899), but the analyses of Schwarz appear to establish the formulæ quoted above.

Esters of Chromic Acid have been prepared by Wienhaus (Ber. 1914, 47, 332) by shaking a tertiary alrohol, dissolved in light petroleum or carbon tetrachloride, with an excess of solid chromium trioxide. The eater remains dissolved or suspended in the solvent, from which it can be obtained by evaporation.

CHROMAMMONIUM COMPOUNDS AND DEPIVATIVES.

chromium compounds then known are classified in his contribution. Prominent authors in the hibliography are Wemer, Pfeiffer, Weinland, and Jörgensen and the following general treatises

havo been puhlished:
Weroer, "Neuero Anschauungen auf dem Cehiete der anorganischen Chemie."
Urbam and Senechal, "Introduction à la

Urbain and Senechai, "Introduction a in chimie des complexes." Weinlund, "Einführung in die Chemie der Komplex-Verbindungen," F. Enke, Stuttgart,

Pfeiffer, "Organische Molekül Verbindungen." Hexammine Chromium Salls (luteo salts), eg. the nitrate, [Cr(NHa)a](HOa)a, are prepared by treating the corresponding purpured salts with ammonia in cold, concentraled solution (Jörgensen, J. pr. Chem. 1882, (2), 23,

229; Mills, Phil. Mag. (4), 35, 245). Pentammine Chromlum Salls (purpureo salts) —Jörgensen (J. pr. Chem 1879, [2], 20, 105) prepared the chloride [CrCl(NH₂)₅]Cl₃ by reducing violet chromic chloride in a stream of pure, dry hydrogen al a red heat, and adding to the chromous chlorido a solution of NH,Cl in strong ammonia. Air was passed through, HCl added, and the muxture boiled, when a carminesaded, and the mitture bound, when a carmin-roloured powder precipitaled. It crystallised in octahedrs, ap gr. 1 687, dissolved in 154 parts of water at 16°, and its analysis corresponded to the formula Ci₂Cr₂(NH₂)₁₀Cl₂ Jorgensen found that only 4 atoms of chlorino are precipitated by silver nitrate, which ronfirms Wemer's formula given above.

Triammine Chromlum Salls -Thesorompounds have been investigated by Werner (Ber. 1910, 43, 2286), who employed triammino chromare, -5, -20), was employed thammino caromium tetracide in their preparation. To a well-cooled solution of 30 g. of chromic acid in 300 c. of water are added 300 c. of pyridine. After temaning balf an hour in a freezing mixture, 750 c. 3% H₂O₂ are added. The precipitated pyridine perchromate is rollected, washed, and added to 90 c.c. of well cooled 25% ammonia. The precipitaled triammine chromium tetroxide CrO,3NH, is rollected after 10 min-utes and washed with waler, alcohol, and ether. Trichtorofriammine Chromium,

Cr{HH,),Cl, results when trammine chromium tetroxide is added to cold, concentrated hydrochloric acid. A bluish grey precipitate is formed, and the filtrate from this deposits the compound after standing 2 days, in dark blue crystals with a greenish tinge. Triaquotriammine-chromium Salts of chromium readily combine with chlords, [Cr(OH)₂(H)₂(H)₃)[Cl, is obtained ammonian or substituted ammonian to form by treating I g, of the dichloroaquotrammine complex rompounds in which the chromium chromium chlorids (contained in the grey atom may be combined with 6 mola of ammonian, precipitate above mentioned) with 8 c.c. of sal, forexample, in hexamium chromium methods: as, for example, in hexammune chromm melhorids with a case of parliam. The [Gr.[HH,1],[C]] The ammona may be substanted by a card groups, the valency and idoda perceptiated by the addition of solid character of the romplex radical being determined polarization and Laus radicals in the molecule. P. Daval has contributed a valuable atudy of It. forms brownish-red hygroscope erystals, the complex compounds of chromaum to Fancia. the complex compounds of thromsum to Pascal's The corresponding bromine compounds have "Traite de Chimie Minérale," tol. x, pp. 1005 been prepared.

Chromi-aquo-triammines have been prepared:

 $[Cr(NH_3)_3Cl_2H_2O]Cl_1$ —blue, $[Cr(NH_3)_3Cl(H_2O)_2]Cl_2$,—reddish-violet, $[Cr(NH_3)_3(H_2O)_3]Cl_3$,—red.

In solution the mono-compound passes into the di- and the di- into the tri-aquotriammino chromitrichloride (Frowein, Z. anorg. Chem. 1920, 110, 107-224).

Schlesinger and Worner (J. Amer. Chem. Soc. 1929, 51, 3520) state that ehloropentammine-chromic ehloride is converted into insoluble chromium trichlorotriammine, Cr(NH₃)₃Cl₃, by heating at 270° in hydrogen ehlorido until the salt turns from palo red to green. This substance reacts slowly with liquid ammonia to yield the original pentammine; it is possibly a polynuclear compound and its properties are quite different from those of the substance of the same composition described by Werner (J.C.S. 1910, 98, [ii], 960). If it is heated at 175° in hydrogen chloride, hygroscopic crystallino ammonium chromic hexachloride,

(NH₄)3[CrCln],

is produced; this substance reacts rapidly with gaseous ammonia with production of the original pentammine. If the triammine salt is heated at 270° in hydrogen chloride a salt having the composition 2CrCl₃,3NH₄Cl is formed, and is further slowly decomposed to anhydrous chromic chloride.

Relnecke's Salt, [Cr(NH₃)₂(SCN)₄]NH₄, tetrathioeyanatodiamminoehromiammonium (Annalen, 1863, 126, 113; Z. anorg. Chem. 1893, 1, 135). To three parts of NH₄SCN in incipient fusion, one part of (NH₄)₂Cr₂O₇ is added in small portions. The red mass is washed theroughly with cold water and recrystallised from water or alcohol. Its solution gives precipitates with the heavy metals and organic bases, alkaloids yielding characteristic crystalline compounds—reineckates (Rosenthaler, Arch. Pharm. 1927, 265, 319. For a modern preparation v. Dakin, Org. Syntheses, 1935, 15, 74).

Chromic Oxalates.—There is an extensive literature of these compounds, their double salts and isomerides (Beilstein, 4th ed. 11, 522, 523, Erg. 11, 226, 227). Special interest attaches to the blue potassium trioxalatochromiato or Gregory's salt, K₃[Cr(C₂O₄)₃],3H₂O, which A. Werner resolved into d- and l- forms by crystallising the strychine salt (Ber. 1912, 45, 3061). The active isomers possess very high rotation and rotation dispersion. Jacger (Rec. trav. chim. 1919, 38, 171, "Optical Activity," New York, 1930) found for the I-salt, [a] for λ 4740, -486° ; λ 5,700, 0° ; λ 5800,+ 515. Potassium dioxalatodiaquoehromiate or ('roft's red salt, trans. K[Cr(C2O4)2(H2O)2]. 3H2O is obtained from hot very concentrated solutions of K.Cr.O, and C.H.O.. The cis isomeride is prepared by moistening a mixture of the powdered substances. With aqueous NH, the cir. salts become green, whilst trans- salts turn brown (A. Werner et al., Annalen, 1914, 406, 261).

For Detection and Estimation of Chromium, see Chemical Analysis.

G. S. B. and J. N. G.

CHROMOFORM is an orange-red crystalline powder formed by the union of methyl dichromate and hexamethylenetetramine. Slightly soluble in alcohol and cold water; readily soluble in hot water. The aqueous solution on boiling, or by treatment with neids or alkalis, evolves formaldehyde.

CHROMOGEN AND CHROMO-PHORE v. COLOUR AND CHEMICAL CONSTITU-

CHROMONE, or benzo-y-pyrone:

is obtained by the action of sulphuric acid on phenoxyfumaric acid (I), whereby benzo-y-pyrone-2-carboxylic acid (II) is produced. This, when heated in vacuo, yields benzo-y-pyrono (Ruhemann and Stapleton, J.C.S. 1900, 77, 1179; Gomberg and Cone, Annalen, 1910, 376, 228).

Chromone crystallises in colourless needles, m.p. 59°, and its yellow solution in cold sulphuric acid possesses a blue fluorescence.

CHROMOTROPE or CHROMO-TROPIC ACID. 1:8-Dihydroxynaphthalene-

3:6-disulpbonie acid.

CHRYSANILINE v. Acridine Dyestuffs. CHRYSANTHEMIN. A monoglueoside of cyanidin (q.v.) isolated from the deep red flowers of Chrysanthemum indicum Linn. by Willstätter and Bolton (Annalen, 1916, 412, 136). The chloride C₂₁H₂₁O₁₁Cl·1½H₂O separated in red-violet leaflets. Hydrolysis of chrysanthemin yielded cyanidin (1 mol.) and glucose (1 mol.). The pierate, difficultly soluble in water, sintered at 165° (decomp.). In its colour renctions-blue-violet with sodium embonate, pure blue with caustic soda, violet-red with sodium acetate, and purplo with ferric chlorideand its high monoglyeosidie distribution between amyl alcohol and aqueous acid, it closely resembles asterin (Willstätter and Bolton, Annalen, 1916, 412, 1149) from the purple-red aster, with which Robinson and Willstätter (Ber. 1928, 61, 2503) have shown it to be identical. That the differences in erystalline form and solubility between these pigments are due to slight impurities is suggested by Murakami, Robertson and Robinson (J.C.S. 1931, 2665) who have synthesised 3-β-glucosidoxy-5:7:3':4'-tetrahydroxyflavylinm ehloride

and shown that it is identical with the chloride of sunder this name in the British and United States chrysanthemin and of asterin and with the Pharmacoprias, and is used in shin affections. product of partial hydrolysis of the cyanidin Chrysarobin was originally thought to be a 3-diglucoside, mekocyanin (Willstätter and Wed, Annalen, 1916, 412, 231) Chrysanthemin chloride is the serond of the remarkable series of anthocyan glurosides which have now been synthesised by Robinson and his collaborators

(see Callisterhin). Chrysanththemin closely resembles the corresponding evanidin 3 galactoside, sdaein, from cranberries both in colour reactions and absorption curves, but differs from it in distribution number (Grove and Robinson, J.CS.

1931, 2723). Pigments identical with chrysanthemin have been isolated from mulberry fruit by Yamamoto (J. Agric. Chem. Soc. Japan, 1934, 10, 1046), from the seed roat of Glycine Soja Benth. ("Kuromame") by Chika Knroda and Mizu Wads (Proc. Imp. Acad, Tokyo, 1935, 11, 189) and from purple-husled marze by Sando, Milner and Sherman (J. Biol. Chem. 1935, 109, 203). In the last case its occurrence together with a 3 glurosids of the closely related flavonol quercetin is quoted in support of the theory of flavonols as the precursors of the anthocyanins It must be noted, however, that cyangdin and quercetin glyrosides are the most commonly occurring types of anthocyanin and anthoxanthin found in nature, so that their occurrence together need not be significant.

R. and G M. Rohmson have identified anthocyanina isomeric or identical with chrysanthemin in a great number of other flowers and trust by means of their qualitative tests (Biochem. J. 1031, 25, 1687; 1932, 26, 1647, 1934, 28, 1712).

Genetical investigations with the Chinese aster by Wit (Dissertation, Gröningen, 1936) show that while pigmentation by cyanidin monoglucoside is sometimes recessive to that by the corresponding 3 5-diglucoside of evanidin. the eyanidin type of pigmentation is dominant to the pelargonidin type and recessive to the delphinidin type, the production of the differently oxidised anthocyanina probably depending upon the action of multiple allelomorphic cenes (see ANTHOCYANS).

R. S.-M. CHRYSAROBIN (B.P., U.S.P.) is a yellow colour from primrose-yellow to bronza and purple, and rollects, possibly as the result of (Fam. Leguminose), a tree inhabiting the forests of Baha (Aguiar, Pharm. J. 1880 [ni], 10, 42; cf Greenish, 151d. 814). The drug was formerly exported by the Portogueso from Gos, but its origin for a long time remained secret, Goa powder is usually mixed with woody frag-By aometimes termed crude chrysarohin extraction of Goa powder with organic solvents $C_{11}H_{12}O_2AC_r$, m.p. 100°, and a triacety] com-(benzene, chloroform) there is left behind on pound $C_{12}H_1O_2AC_r$, m.p. 238°. evaporation a yellow crystallium powder, introduced in 1875 by a Bombay firm under the was not solated by Tutin and Clewer as such,

pure, or nearly pure, substance; in reality it is a complex mixture, consisting chiefly of reduced di- and tri-hydroxymethylanthraquinones, which erestalliso together and are difficult to separate by purely physical means (cf. liesse, Annalen, 1899, 309, 53). Later the namo ehrysarohin was applied to one of the pure constituents, but it is better to reserve this name for the commercial mixture. Tutin and Clewer (J.C.S. 1912, 101, 290) found the

following perrentage composition for three samples of commercial chrysarobin: 1. Chrysophanic acid (=chrysophanol) 2. Emodin monomethylether1 3. Emoden . trace

4. Cruds anthrenol of chryso. phanie and

5. Anthranol of emodin monomethyl ether . small amounts 6. Monomethyl ether of de-

hydroemodinanthranol 411 134 7. Ararobmol 8. Inseparable mixture

partly amorphous aubstances. About 3 or 4 parts of the former to 1 of the latter

Of these No 1 is a methyldihydroxyanthraquinone, No. 3 is a methyltrihydroxyanthra-quinone, Nos. 4 and 6 result from Nos 1 and 2 respectively, by reduction ; they are derivatives of anthranol (9 hydroxyanthracene). No 6 is a No. 7, of less strongly reduced enthraquinone unknown constitution, is also an anthraquinone

denvative, related to No. 1. Chrysophanic acid anthranol, C. H., O., has the constitution

and was called chrysarobin by Jonett and Potter (J C.S. 1902, 81, 1578). It is obtained from commercial chrysarobin by extraction with bight petroleum; by ahaling its chloroform solution with aodium carbonata trares emodin are removed, and after evaporation crystalline mixture of substances obtained from of the coloroform the residue is crystallised from Araroba or Gos powder. The latter varies in much ethyl aretate, when it forms lemonmuch ethyl aretate, when it forms lemon-yellow scales, in p. 202° (rorr.) (regenerated from its acetyl derivative the aubstaoro melts oxidation of the resin, in the cavities of the at 204°). It is insoluble in sodium carbonate, stems and branches of Andira Araroba Aguiar soluble in caustic alkalis to a yellow solution, which becomes red by passing air through it, whereby it is oxidised to chry sophanic acid. Tha change is complete in 2 hours, as shown by measurement of the oxygen absorbed (Tutin and Clewer, ie.). The reverse changa is brought about by hydriodic acid at 130°-140°. Chrysophame ments from which it is freed by afting; it is acid anthranol dissolves in sulphuric acid with a yellow colour; it forms a diaretyl rompound

name Chrystrobin, which has since been included but its presence was inferred from aerial oxida-

tion experiments, which converted it into emodin]

monomethyl ether.

The monomethyl ether of dehydroemodinanthranol, C16H12O4, is characteristic of cbrysarobin. According to Eder (v. infra), its constitution is most probably:

Tutin and Clewer converted it by boiling hydriedic acid into emodin anthranol, a substance containing a methyl group less and 2 hydrogen atoms more; by oxidation with chromic acid they converted it into emodin

methyl ether.

Of the substances 1-6 chrysophanic acid had also been found by Attfield, its anthranol by Jowett and Potter, and emodin monomethylether by Ocsterle and Johann (Arcb. Pharm. 1910, 248, 476), but this was the full extent of the agreement among the various investigators of chrysarobin until Eder (Arch. Pharm. 1915, 253, 1; 1916, 254, 1) confirmed all the principal results of Tutin and Clewer. Ho oxidised tho commercial substance with air in alkaline solution, and considers that hydrogen peroxido is formed in this reaction, and is responsible for further changes. Ho thus obtained the following percentages: emodin 0·2, emodin methyletber 9·3, chrysophanic acid 22·7, dchydro-emodin anthranol monomethyl ether 18 (=chrysarobol of Hesse?), amorphous products 26. It should be remembered that most of the chrysophanic acid and some of the emodin methyl ether were originally present as anthranols. In his second paper, Eder acetylated and benzoylated chrysarobin and obtained by subsequent hydrolysis up to 14% of chrysephanic acid anthranol, up to 9% of emodin anthranol monomethyl ether, and up to 4% of emodin monomethyl ether, but no chrysophanic acid. The chief discrepancy is that Eder did not find the last of Tutin and Clewer's substances, ararobinol. This, according to the latter authors, has the composition $C_{22}H_{16}O_5$, and forms yellow crystals, decomposing without melting at about 225°. It dissolves in alkali with a yellow colour, but, like the methyl other of dehydroemodin anthranol, and unlike chrysophanic acid anthranol, it is not readily oxidised by air. It gives a delicate and characteristic reaction with sulphuric acid; a mere trace forms an orange solution, which on gentle agitation gives intense blue streaks; the solution then becomes green, and finally dull grey. Ararobinol gives a triacetyl derivative (decomp. at 225°), is reduced by hydriodic acid to dehydroararobinol, and oxidised, but not quantitatively, by cbromic acid to chrysophanie acid.

nally as a purgative. Probably its chief constituent, chrysophanic acid anthranol, does not phanic acid), as is the case with aloin (Meyer and Gottlieb, "Experimentelle Pharmakologie," DYESTUFFS.

Berlin and Vienna, 1933, p. 240). The chief use of chrysarobin is, however, as an ointment in skin diseases (psoriasis), and the active constituent appears to be the anthranol of chrysophanic acid. According to Unna's experiments (Baudisch, Ber. 1916, 49, 179), the only active synthetic substitutes are the anthranols of and of 1:8-dihydroxy-antbra-1-hydroxy- and of 1:8-quinone. The latter, cignolin,

is more active in psoriasis than the anthranol of chrysophanic acid (chrysarobin), because it bas not the dystherapeutic methyl group.

Baudisch explains the effect of the 1:8bydroxyls in accordance with Pfeiffer's views by

the formula:

"Dermol" is prepared by precipitating a bismuth nitrate solution by a solution of chrysophanie acid in sodium hydroxide. 'Anthrarobin" is the anthranol of alizarin.

The different methods of extraction cause considerable variation in the composition of commercial chrysarobin and in its physiological action (Pharm. J. 1911, [iv], 87, 630). Henco Tutin and Clewer consider it fallacious to ask for compliance with exact requirements. (The United States Pharmacopæia states that 1 g. of chrysarobin dissolves in 385 c.c. of alcobol, 13 c.c. of cbloroform, 160 c.e. of other, 30 c.c. of benzene, and 180 c.c. of earbon bisul-Pure chrysarobin dissolves in phide at 25°.) concentrated sulphuric acid with a yellow colour, and is insoluble in dilute sedium hydroxide; concentrated sodium bydroxide dissolves it with a green colour. Chrysophanic acid, on the other hand, yields a red solution with concentrated sulphuric acid and dilute sodium hydroxide (Liebermann and Seidler, Ber. 1878, 11, 1603). Another method to distinguish chrysarobin from chrysophanic acid is to mix 0.05 g. with 0.2-0.3 g. sodium peroxide in a porcelain capsule, add 5 e.e. of alcohol, and after 5 minutes 15 c.c. of water. Chrysarobin yields a wine-lees colour, not destroyed by dilution with water, changing to yellow on the addition of acetic acid; chrysophanic acid yields a cherry red colour, becoming brighter on Chrysarobin is occasonally employed inter- dilution (Piñerûn-Alvarez, Ann. Chim. Analyt. 1907, 12, 9).

exert a purgative action until it has been chryso-derivative of esculetin (6:7-dihydroxycoumarin). CHRYSAZIN v. ALIZARIN AND ALLIED The highest boiling fractions of coal tar are the richest source of chrysene (for typical treatments or pitch, see C. Zerbe and F. Eckert. Brennstoff.
Chem. 1934, 15, 28; J. W. Cool. and C. L.
Hewett, J. C.S. 1933, 3953, sithough it occurs in
tars from many natural products. It is formed
by simple distillation of some natural fats, oils and resins, and by drastic debydrogenation of many natural compounds possessing a related ring structure, eg cholie acid (L. Ruzicka and co-workers, Helv. Chim Acta, 1934, 17, 200), toad poisons (H. Wieland and G. Hesse, Annalen. 1935, 517, 22), follicular hormone (A. Butenandi and H. Thompson, Ber. 1934, 67, [B] 140). Chrysene anses also as a result of pyrolytic treatment of many simpler compounds, eg phenol (A Hagemann, Z angew Chem. 1929, 42, 355), tetrabn (N. A. Orlow and N. D. Lichatschew, Ber. 1930, 63, [B], 2179), natural gas (S. F. Buch and E. N. Hague, Ind. Eng. Chem. 1934, 28, 1009). The sction Eng. Chem. 1934, 28, 1009). The sction of caustic potash at 300°-310° on indene yields a little chrysene (R. Weissgerber and C. Seidler, Ber. 1927, 60, [B], 2088), and considerable yields pletinum cetelyst (N. Orlow and M. Belopolsky, Ber 1929, 62, [B], 1226).

The need for synthetic approaches to condensed eyelic systems of the steroloid type has resulted in the elaboration of several syntheses of chrysene and neer derivatives. Thus cyclo-hexanone and R(1 naphthyl) thyl magnesium bromide yield a tertiary alcohol which is cyclosed to a hydrochrysene and then dehydrogenated with selenium (J W. Cook and C L. Hewett, JCS 1933, 1093, 1934, 365) Variations using Δ'-cyclohexenyl magnesium bromide (J W Cook and A. Dansi, JCS. 1935, 500) or & phenyl ethyl magnesium bromide (L. Ruzicka and H. Hoek, Helv. Chim. Acta, 1934. 17, 470) with a tetralone have also been effected Again, an amorphous tetrahydrochrysene has been synthesised by cyclising \$ (2 phenanthryl) propionic acid and reducing the resulting ketotetrahydrochrysene by the Aemmensen method (R. D. Haworth end C. R. Mavin, J.CS. 1933, 1012). A third regent synthesis of the chrysene ring system due to J von Braun and G. Irmisch (Ber. 1931, 64, 2461) and extended by R Robinson and G Ramage (Nature, 1933, 131, 205; J.C S 1933, 607) consists essentially in a double eyclisation of \$\$ diphenyl adipic send or its chloride to a mixture of isomeric diketohexahydrochrysenes

Removal of other hydrocarbons from the chrisene fraction of coal tar with carbon dualf lade followed by further purification by usual means yields chrysene which is persistently contaminated with a yellow importly resisting reparation by crystallisation, but which may be

removed by oxidiasing egents, eg perchloric acidi.

K. A. Hoffmann, Ber. 1910, 34, 1059, natire
sed and alcohol (Liebermann, Annalen,
1871, 155, 299); see also Schmidt, J. pr. Chem.
1874, [ul. 9, 250, 270). Other methods recommended for the purification of chrysene are
separation through the characteristic 2.7
diastroanthraquinone complex (Schmidt, Ic.)
and combination of the impurity with males
ashydride (E. Clar and L. Lombardt, Ber. 1932,
25, [Bl. 1411] For small quantities the method
of chromatographic adsorption (r. Christical,
ANALYSIS, Vol. 11, p. 628) is mutable, A. Winter
attern and co-workers having identified the
continuous (1934, 22, 23, 24, 24), and Chem.
1934, 250, 146, 156).

T. Physiol. Chem.
1934, 220, 246, 156).

Pure chrysene forms colourless rhombic plates mp 250°, bp 448°, which exhibit in the solid state or in solution a red violet fluorescence it is best characterised by the 27-dinitro anthraquimone complex (Schinidi, Ic., Haworth and Mavin, Ic.). For a colour reaction of chrysene, see Ditz (Chem Zig, 1907, 31, 445)

An J., Linhafetenew, Rev. 1304, S. J. 1935, 1943, Substitution products of chrysen ere obtain tags. Clem. 1934, 26, 1003. The extion and the product of the

719, 1278)
Chryseno has not yet found extensive industrial use, although patents have been secured for true use in the manufacture of dystalfs. Thus 1.4 quamons are condensed with an ambight boiling solvents and in presence of mild oxidising agents to yield pigments or, on further sulphonation, dystalfis (FP, 754987). Again, chrysene is districted, reduced to diammochysene, end then subjected to a double Skrapp synthesis to a condense of the control of

A И С.

CHRYSIN. 5.7 dibydroxyflavone,

119 CHUFA.

is obtained from poplar buds (Piecard, Ber. 1874, 7, 88; and Darier, ibid. 1894, 27, 21). It is a member of the quereitin series (Perkin, J.C.S. 1896, 69, 1443), and dyes cloth mordanted with aluminium a pure yellow; with chronium, an orange-yellow; and with iron, a pale chocolate (Perkin, ibid. 1897, 71, 818).

CHRYSOBERYL. Aluminate of beryllium, BeAl₂O₄, crystallising in the orthorhombic system, and used as a gem-stone. It is usually met with as well-formed crystals, which as a result of twinning often simulate hexagonal forms, or as rolled pehbles in river gravels. Sp.gr. 3.65-3.75; hardness 81, taking the third place amongst minerals, being oxceeded only by diamond and corundum. Few minerals are so resistant to chemical reagents as chrysoberyl; it is not attacked by acids nor fusible before the blowpipe, and is decomposed only by fused alkalis. The colour ranges in various slindes of yellow and green. The variety alexandrite is emerald-green in daylight, but violet-red in lamplight. Another gem variety, known as cymophane or "oriental eat's-eye," shows a bluish milky band of reflected light which travels across the convex polished surface as the stone is moved about. Chrysoberyl occurs embedded in granite and erystallino sehists. Material of gem quality comes from Ceylon, Brazil and the Urals. Colourless crystals have been found in the ruby mines in Upper Burma and in the Gold Coast in West Africa. By reason of its high degree of hardness it has occasionally been used for jewelling watchbearings.

L. J. S. CHRYSOCOLLA. A hydrated silicate of copper, sometimes of importance as an ore of copper. The formula is usually given as CuSIO2,2H2O, but the composition is variable, the mineral often being mixed with various impurities-silica, alumina, copper carbonate or phosphate, limonite, etc. On these differences in composition several trivial varieties have been distinguished by special names. The mineral is of secondary origin, occurring in the upper portions of deposits of copper ores. It forms encrusting or botryoidal masses, which may be vitreous and translucent, resembling opal or enamel in appearance, or dull and earthy. The colour ranges from bright green to bright blue, and there is a wide range in the sp.gr., 2.0-2.8. Under the microscope it sometimes shows a minutely fibrous structure, and small acicular crystals from Idalio have been described, but with variable and uncertain optical data and sp.gr. 2.4 (J. B. Umpleby, 1914). Interbanded with the microerystalline material is often some optically isotropic material, and for this amorphous equivalent of chrysocolla the name cornuite has been proposed (A. F. Rogers, 1917).

A variety known as pitchy copper-ore (Ger. Kupferpecherz), dark brown or black in colour and with the appearance of pitch, is of some importance as an ore of copper, particularly in Chile. It consists of a mixture of chrysocolla and hmonite (F. N. Guild, Amer. Min. 1929, 14,

Belgian Congo, Chile, Arizona, Idaho, etc. The name chrysocolla, from χρυσός, gold, and κόλλα, glue, was applied by the ancients to materials used for soldering gold, and included, besides certain copper minerals, also

CHRYSOGEN. The crude orange-yellow hydrocarbon responsible for the yellow colour and green-yellow ultraviolet fluorescence of crude anthracene. Capper and Marsh (J. Amer. Chem. Soc. 1925, 47, 2847; J.C.S. 1926, 724) studying absorption and fluorescence spectra, concluded that chrysogon was either naphthacene (I), or 1:2-benzanthracene (II) (see also Cook and co-workers, Proc. Roy. Soc. 1932, B, 111, 469; Morgan, J.S.C.I. 1928, 47, 132).

Winterstein, Schon and Vetter (Z. physiol. Chem. 1934, 230, 158) separated chrysogen from anthracene by chromatographic analysis and completely identified it with naphtha-

CHRYSOLITE, a clear, transparent, yellowish green variety of olivine (MgFe)2SiO4.

CHRYSONAPHTHAZINE v. Azines. CHRYSOPHANIC ACID (Chrysophanol) v. CHRYSAEOBIN.

CHRYSOPRASE v. CHALCEDONY.

CHRYSOTILE v. Assestos and Ser-PENTINE.

CHRYSOTOLUAZINE v. AZINES.

CHUFA. Earth almond. The tubers of Cyperus esculentus Linn., grown extensively in central and southern areas of the United States. The tubers, which are 0.5-1.0 in. in length, have a nut-like flavour and are said to resemble the water chestnut. They are sometimes roasted and used as a substituto for coffee. The plant is also valued as forage for pigs. The fat content of the tubers averages 20-29%, starch 12-30%, and sucrose 15% (Power and Chesnut, J. Agric. Res. 1923, 26, 69; Pieraerts, Mnt. Grascos, 1924, 16, 6674; Anon. Giorn. Chim. 1933, 27, 297).

Baughman and Jamieson (J. Agric. Res. 1923, 26, 77) record a detailed examination of the oil and give the following characteristics: d15 0.918, 313). Chrysocolla of the more ordinary type refractive index 1.4662 (25°), saponification has been mined for copper in South Australia, value 191.3, iodine value 76.89, Reichert-Meissl 120 CHUFA.

Lignoceric acid			0-3
Arachidic acid			0-5
Steame acid .			52
Palmitic acid .			11-8
Myristic acid			trace
Oleic acid			733
Linolic acid .			59
Unsaponifiable	acid		30

A phytosterol, m p. 134°-135°, was also found-A related species, C. rolundus, from Japan. yields 1% of an essential oil which, according to Kimura and Ohtani, contains 32% of cyperene (C11H24) and 49% of experol (C11H40).

CH'U-MA & CHINA GRASS.

CHURCHITE. A hydrated phosphata of cerium (Ce,O, 51 87%) and calcium, occurring as grey fan-like aggregates of acicular crystals encrusting quartz and elate in a Cornish copper lode. Unfortunately nothing is known of the exact lorality of the mineral.

L J. S.

CHYMASE a synonym for REVEET. CHYMOTRYPSIN. Anactive protectlytic enzyme obtained by the action of a minute quantity of trypsin on a crystalline peotein contained in acid extracts of pancress (Kumitz and Northrop, J. Gen. Physiol. 1935, 18, 433). This enzyme hydrolysce casein more slowly than trypsin but carries the hydrolysis further and attacks different bakages.

E. F. A.

CIBACET 1. ACETATE SILE DYES. CIBANONE DYES. Blue G, 3G, RA; Black B ; Brawn B; Ohve B ; Grange R. CR ; Red 4B ; Violet BW ; Yellow R (r. ANTHRAQUIVONE DYESTUTES

CICHORIIN, C1, H100, 2Aq, m.p. 213°-215°, [a]p -1045°, is the glycoside present in Gickorium Intigous, chicory.

It is isomerie with asculin differing only in

the Position of the sugar residue According to K. W. Merz (Arch. pharm 1932, 270, 476), it is hydrolysed to glucose and cichorigenin, C.H.O., which is 6.7 dihydroxycomarys. In eichorum the glucose residue is attached to position 7, whereas in asculm the attach-

When eichorun is first methylated and then bydrolysed, 6-methoxy-7-hydraxycoumarin, s.e.

ment is at position 6. ecopoletin, is obtained.

E. F. A. the possonous

CICUTOXIN, C19H24Op the poisonous principle in water-hemlock (Cicuta tugana) is contained in the tubers to the extent of 0-3-0 4%. It is found also in C. virosa. It is a yellow resin-like substanco which readily decomposes and polymerista above 50°, it has des 0 9659 and not 1-5885. Unites energetically associated with beverages conforming to certain with bromine and ignitra spontaneously in specified legal standards. Other countries vary contact with strong nitre acid. On dry distilla-in the settent and methods of protection and in tion it yields an oil, C₁H₂₁O₂, of a²⁰ 0306, some they are practically non existent. Thus und Dattilled in a current of steam it gives an oil, der the general name of caler there are to be found C₁H₂₂O₂ of a pleasant odour and which beverages of an extremely wide range of types.

value 0-2, Polenske value 0-3, acetyl value 4 55, | gradually polymerisea to a transparent semiacid value 1-70. The calculated composition sold rean. On oxidation with nifric acid circulated fatty acids is: gen cyanide, isobutyrio acid, and acetyl-2-cyclopentanone, bp. 73 5 15 mm. Compounds of cieutoxin with lead, barnum, hydrogen chloride, ammonia, bramine and iodine are known. Cieutoxin appears to be a derivative of 4 pyrone (Jacobson, J. Amer. Chem. Soc. 1915, \$7, 916); it exhibits absorption bands at λ 570-64 and λ 650-680 (Švagr, Chem. Listy,

1923, 17, 166). CIGNOLIN, the anthranol of 1.8 dibydroxy.

anthraquinone (v. CHRYSAROBIN).

CIDER or CYOER is a beverage produced from the juice of apples by alcoholic fermentation. A corresponding beverage, perry, is made from the suce of pears; this so closely resembles eider in general type and qualities, methods of production, and centres of manufacture, that for the purpose of this article the two drinks can be treated in common and distinctive reference only made to perry when occasion requires.

Both ara beverages of considerable antiquity, the earliest extent mention of cider in this country relating to its production in Norfelk as far back as the thirteenth century. In Franco it was being made in the sixth century, and probably still earlier. That country and Great Britain era the two leading producing Great Driam era the two leading producing countries at the present time, the output of the former in some years exceeding 400,000,000 splices and being many times greater than that of the latter. The approximate area of the French cider finit orchards, centred mainly in Normandy and Brittany, is 1,330,000 acres, and of the English 55,000 acres, The latter are situated principally in the counties of Devon, Hereford and Somerset, with Gloucester and Worcester next in order, Cider fruit and cider production also extend into Doract, Monmouth and other counties adjacent to those named. In Norfolk and, to a less extent, Kent, eider is also made commercially, but in those districts cider varieties of apples are not grown on any considerable scale and "market" (i.e. eating) apples are used instead

Cider in some form is made in most other countries in which apples flourish, but in their case for the most part vintage varieties of the fruit are not grown. In Europe Germany, Switzerland and Spain particularly have gamed some repute for their cider. The beverage produced in the United States and their cider. The Canada in some quantity under this name has been during recent years mostly unfermented apple juico, although latterly in Canada efforts to develop the manufacture of the fermented

articla have been vigorously pursued.

The initial definition of cider given above indicates a beverage with a material content of alcohol, and in Franco in particular the use of the name is pratected by law and must only be

differing in respect of original juice content, chemical composition, and form of presentation. In the matter of juice content France, for example, enforces the application of specific forms of name designation indicating (a) whether the article is produced from undiluted or diluted juice and, if the latter, (b) the degree of dilution.

In England no legal standards have been formulated, hut under a regulation of the Ministry of Health, which came into force on January 1, 1927, no prescrvative except sulphur dioxide is permitted. The maximum content of sulphur dioxide allowed is 0.02%, which is equivalent to 14 grains per gallon. The introduction of a National Mark Schome for Cider in 1931 by the Ministry of Agriculture and Fisheries has lcd, however, to the establishment of a voluntary standard for English ciders which must be attained before the ciders are entitled to hear the National Mark. According to this standard, the cider must be made from apples grown in England and Walcs. It must be made from cither the pure juice of apples or a diluted juice centaining not more than 25% of its own volume of a syrup of pure cane or pure heet sugar. The original specific gravity of the juice must not be below 1.040 and the percentage of volatile acid (as acctic) must not exceed

The chemical composition of eider in relation to the centents of alcohol, sugars, acid, and tannin is very variable and is responsible for a series of types of the heverage differing widely in taste and physiological action. Total sugar content determines the classification under the heads of sweet, medium, dry and extra dry, the percentage ranging from 6-7% hy weight or over in the case of sweet eiders to helow 1% in the case of the extra dry types. Alcoholic strength tends to vary in the opposito direction and may be but little over 2% hy weight in sweet ciders and exceeding 5-6% in dry and extra dry: in special cases, where a higher content is required, this generally cannot be obtained directly from untreated pure juices on account of their limited initial sugar content and is provided either through the fermentation of added sugar or by direct fortification with spirit. The latter methods make it possible to secure a high alcohol content, even in the sweeter types. Acid and tannin determine the degree of acidity, and astringency and hitterness, respectively in the flavour. Ciders containing over 0.4% of malie acid rank as brisk ciders and the low-acid types generally range in acidity between 0.2 and 0.4%. A tannin content in excess of 0.2% communicates a definite "hittersweet" character to the flavour and is generally accompanied by at least a fair degree of fullness in body; below that figure this "bittersweet" character is normally not marked on the palate. Ciders of a low-acid, somowhat bittersweet character are generally favoured in France; in Great Britain a brisker, less bittersweet type is in most demand except in the eider-making country districts, where a rougher and more bitter, dry type suits the local taste.

Cider is supplied either in eask or bottle.

nearly so, any slight sparkling condition being generally the result of a slow after-fermentation. In hottle, sparkling cider is the rule, the carhon dioxide to which the "life" is due being seenred mostly by natural conditioning in the hottle or hy artificial carbonation prior to hottling. The latter method predominates in commercial production. A third type of sparkling eider is that produced by the true "champagne" method, the liquor heing first strongly fermonted in bottle and then disgorged to remove all sedi-

Although a heverage coming within the ordinary definition of cider can be made from any kind of apple, the result is likely to he inferior and lacking in commercial appeal nnless either the fruit is composed of vintage apples exclusively or of a mixture in which vintage fruit is the predominant constituent, or, alternatively, special methods of treatment are adopted. Fruit falling within the class of "cating" apples, i.e. dessert and culinary varieties, is characterised by a deficiency of tannin and, usually, an excess of acidity from the vintage point of view. Cider made therefrom by usual procedure is typically thin, lacking in hody and character, more or less oversharp (according to the variety), poor-coloured, and coarse-flavoured as a result of an excessive rate of fermentation of the juice.

Since in practice apples unsuited for dessert or culinary purposes are normally sent to the cider mills, they tend undeservedly to be ranked as vintago fruit. Strictly, that term should he reserved for varieties yielding a juice which hy fermentation and after-treatment will produce a cider of good quality in its particular class, irrespective of any blending requirements. Quality in this connection includes characters such as aroma, flavour (in respect of fruit character as distinct from the aspects due purely to sugars, acids, and tannins), body and low nitrogen content. The sum total of such characters may he expressed hy the term "vintage quality." A cider apple proper may, therefore, he defined as one normally possessing vintage quality of a satisfactory

The individual varieties of cider apples show a very wide range of composition in respect of constituents which are of vintage importance. Sugars, for example, may range from 7 to over 20%, malic acid from 0.1 to over 1.5%, and tannin from 0.1 to over 0.6%. The specific gravity of the juice ranges generally hetween 1.035 and 1.090. Although not an exact index of the total sugar content of the juice owing to the presence in the juice of other soluble solids. the specific gravity affords a very fair approximation, by far the greatest proportion of the soluble solids being sugars. A juice with a specific gravity of 1-040 usually contains about 8% of total sugar; of 1-050, approximately 10%; and of 1-060, approximately 14%

Cider apples are classified according to the chemical composition of their juice in respect of the amounts of malic acid and tannin normally present. They fall into three main Draught eider in the wood is usually still or classes—sharp, sweet, and hittersweet.

As a result of the examination of a large of them in the "sweet" class, the palate usually unmber of English cider apples an extended ranks them as mild "bittersweets" Their classification enabling varieties to be grouped so ciders unblended resemble those of Group V. that their blending requirements may be closely indicated has recently been prepared by the Long Ashton Research Station, which eerves as the research centre for the cider

industry in this country. This classification is as follows:

SHARP CLASS.—Group I.—Full Bittersharp (Forwhelp Type). Varieties with juices normally containing not less than 0-750; make acid and

0-18% tannin.

The members of this group are very acid Their ciders are too sharp and in taste. too much inclined to hitterness for use unhlended. Group II .- Full Sharp (Frederick Type)

Varieties with juices normally containing not less than 0-75% make acid and not more than 0-180 tannin

Ciders made from these varieties alone are

too sharp and generally lack body. Geoup III .- Medium Bittersharp (Kingston Black Type) Varieties with juices normally

containing between 0-45 and 0-75% maho acid

and not less than 0-18% tannin
Into this group fall most varieties that yield ulatable and well balanced ciders unblended, Ringston Black being the outstanding example In some instances and in certain seasons, especially when body is lacking or some trait of flavour is abnormally perminent, blending

gives an improved beverage, in such instances they require the addition of an already belanced blend in quantity sufficient to give the character sought for Group 11 - Medium onerp vocality contain-

ing between 045 and 0.750 make acid and

not more than 0-18% tannin. For this group, owing to the low tannin content, hlending with "bittersweet" material is generally essential. Only infrequently car blending be dispensed with Because of the medium acid content the material used for blending needs to be already balanced in respect of acidity.

Ciders made from the varieties of this group unblended, while usually of about the right standard of acidity, generally lack body and aufficient "tannin" character.

SWEET CLASS. Group V .- Sweet (Swee Alford Type). Varieties normally containing less than 0-450 make and and less than 0-182

The members of this group are the "sweet" varieties proper. By themselves they generally yield eiders insipid to the palate because of

deficiencies of acid and tannin. BITTERSWEET CLASS, GOOD VI. - Mile Bittersweet (Belle Norman Type). Varieties normally containing less than 0-45% make seed and between 0-18 and 0-25% tannin-

This group is sufficiently numerous to justify Its group is sumiciently unmerous to justify its apparation from Group V. It includes a superparties of the individual conof variety that is on the border line between the
sarret' and "bitterweet" classes. While
have been obtained by the analyses of three in many seasons chemical analysis places several typical French ciders.

but with rather more character,

Group VII.—Medium Bittersweet (Knotted Kernel Type). Varieties normally containing not more than 0-45% malie acid and appeoxi-

mately 0 25-0 4% tannin

This group is closely described by its title "medium bittersweet," Ciders made from its members unblended have that character very definitely marked as well as distinct lack of acidsty. The tannin percentage varies con-siderably with most varieties according to season-this applies to Geoups VI and VIII also and occasionally places them chemically in the Group VI range; but the palate marks them almost invariably as "medium hitter-

Group VIII.—Full Bittersweet (Strouberry Norman Type). Varieties normally containing not more than 0-45% make acid and not less

than 0-4% tannin.

sweets."

Members of this group are pronounced "bittersweets" with the tannin character generally overwhelming to the palate. That feature distinguishes their ciders from those of the Group VII varieties, which otherwise they closely resemble. Not infrequently this character renders it impossible to use them in hlends

except in low proportions,
While the chemical composition of the apple does not necessarily indicate the value of that apple for eider making, nevertheless vintage quality is closely correlated as a rule with the composition of the juice. Apple juice is a liquid containing many constituents, of which the chief in quantity are the sugars (lievulose, glucose and sucrose), make acid, tannins, pectin and minerals. Typical analyses of three varieties of French apple juices are given in the following table:

1				
3		Bedan	St.Martla	Boutelile
1	Density of purce .	1063	1060	1055
	Water	83 15	84 34	85 75
1	lose	11-10	10 00	9 10
ı		0 28	0.28	0.30
Ş	Niteogen	0.055 0.15	0 062 0 12	0 664 0-13
	Pecture and Gume	0 32	0 51 0 36	0 42
ŗ	Unidentified com- pounds, fats,	[]		
-	aorbite, etc.	3 315	2 948	2 596
3		100 000	100 000	100 000
ł	_	I		

The nature of the ash constituents and the

	Ille-et- Vilaiae	Manche	Seine-Inf.
	%	%	0,000
Silica	0.0017	0.0010	0.0020
Phosphoric acid .	0.0229	0.0205	0.0225
Sulphuric acid .	0.0145	0.0065	0.0225
Chlorine	0.0023	0.0020	0.0225
Iron and alumin-			}
ium oxides	0.0017	0.0015	0.0225
Calcium	0.0050	0.0042	0.0100
Magnesium	0.0037	0.0080	0.0150
Potassium	0.0970	0.0113	0.0730
Sodium	0.0020	0.0021	0.0015
Total ash	1.0805	1.0980	2.0787

Apple tannins have not been very closely studied, but they resemble more the products from coffee, eachon, or quinquina than oak-gall tannin. Their oxidation by means of an oxidase is mainly the cause of the characteristic amber colour of ciders.

The nitrogenous compounds of apple juice consist of complex proteins, acid amides, and traces of ammonia. In freshly expressed juices the nitrogen content usually ranges from 0.15 grown in grass orehards, the type of tree being to 0.03%. During the course of fermentation that known in horticulture as a "standard," a material quantity of the nitrogen content is removed owing to its assimilation by the yeasts and the residual amount in most finished eiders is approximately 0.04%. These constituents have an important bearing on the quality of the eider since they are mainly responsible for the rate of fermentation of the juice. If the nitrogen content is high, the fermentation of the juice is rapid and a coarse type of eider results. A low content of nitrogen is associated with a slow rate of fermentation and a product of superior quality.

The composition of the juice of any individual variety is subject to considerable variation, both from year to year and when obtained from samples of fruit grown under different local conditions. Many factors influence its composition. Among these the more important are climate, soil, condition of ripeness of fruit at the time of expression of the juice, methods of enlture of the tree, age of the tree, the size of the crops borne by the tree during the previous two or three years, and manufal treatment given to the tree.

Perry pears may be distinguished from other varieties of pears by their possession of superior vintage quality, as in the analogous case of the eider apple already referred to. The same general remarks as regards composition and its liability to variation apply in their case too. Owing to their relatively small number, no systematic classification has bitherto been established. The features of any classification, if f rmulated, would probably fellow along very similar lines to those of the apple classification outlined above.

The possession of high vintage quality does not necessarily entitle the variety in question

into account for economic reasons. Thus, for a variety to qualify for inclusion in a selected list, it must attain a suitable standard in respect of both vintage and orchard quality. The important characters concerned with the orehard quality of a .ariety are the capacity of the tree to grow adequately and to maintain a reasonable state of good licalth over a long period of years under grass orchard conditions, freedom from serious susceptibility to damage from insect pests and fungus diseases, good eropping quality, and individual fruits of at least fair size.

During the past quarter of a century both the vintage and orehard qualities of many varieties of eider apples have been investigated and, as a result, the list on page 124 of approved sorts prepared by the Long Ashton Research Station. Some of the varieties included in the list, such as Kingston Black, Foxwhelp, and Cap of Liberty, need to be grown under suitable local conditions and on appropriato soil, but most of the varieties named can be recommended where local conditions favourable for apple culture in general.

a form with a single main stem at least 6 ft. in length from ground level to the beginning of tho head of the tree. This type of tree is necessary to enable use to be made of the grass sward of the orchards for grazing by livestock. Cultiva-tion in grass is preferred to cultivation in tilled land on account of reduced costs of cultivation, the better harvesting conditions for the fruit and the lower nitrogen content obtained in the juice. The individual trees should stand a minimum of 30 ft. apart and many strongly growing varieties need a distance of 35 to 40 ft. apart. An adequate and economic control of pests and diseases can be obtained by spraying in alternate years with an egg-killing winter wash and a spring spraying with line-sulphur respectively. The sward of the orehard should bo kept close by grazing. After the heads of the trees have been suitably shaped and established in their earliest years little pruning is required beyond an annual overhaul to remove any interlacing or dying and dead branches. Orchard management along these lines is now being more and more extensively practised, but the older generation of orehards in the West of England affords many examples of the deplorable results of neglect of proper management.

The earliest-ripening varieties of eider apples begin to ripen in September and the latest aro not fit to gather until the early winter. The correct time for harvesting needs to be decided for each variety separately; it is when the apples fall freely when the tree is lightly shaken. Premature harvesting means a loss of quality in the ultimate eider, since the fruit does not acquire its full vintage quality until it has ripened to a comparatively mature state on the tree. Normally the fruit at this stage of maturity has not yet reached its optimum condition for eider-making and needs to be to a place in a selected list of varieties to be maturity has not yet reached its optimism recommended to growers for culture. What condition for eider-making and needs to be may be comprehensively termed the "orchard stored for a period before that point of condition of the conversion of qualities" of the variety have also to be taken is reached as a result of the conversion of

	Name of vintage class of variety.					
flarvesting season.	Sharp.	Sweet.	Bittersweet.			
Early (to mid October)	Backwell Red	Court Royal	Belle Norman (syn Perthyres) Major Royal Jersey II White Jersey White Norman			
Mid-season (mid- October to early November)	Cap of Liberty (sym. Red Soldier) Dymock Red Soldier) Dymock Red Devon Dourhelp Frederick Kingston Rlack Langworthy (sym. Sour Natural, Wyatt'a Seedling) Porter'a Ferfection Rejuvenated Foxwhelp Stoke Red Yellow Styre	Berkeley Pippin Eggleton Styre Sweet Alfard Sweet Coppin Woodbase II (syn. Northwood, Sweet Woodbase)	Bulmer'a Norman Ellis' Bitter Knotted Keroel Masters' Jersey (syn. Harry Masters' Jersey) Royal Wilding Tremlett's Bitter Yarlington Mill Jersey			
Late (November and later)	Crimson King Lambrook Pippin Reinette Ohry	Woodhine I (1978 Slack-ma Girdle)	Dahnett Dove Modaills d'Or I Michelm Reine de Pommes Silver Cup Tardive Forestier.			

the best results are to be obtained.

thorough.

For the expression of the juice considerable that the drainage racks above and below improvements have been introduced in recent each are nearly in contact. years in the nature and construction of the lit is usual to press the pomace immediately mills and presses and the accessory equipment. after delivery from the mill, but many old

starch to sugars and various changes in the Most modern cider mills are of the grater type-pectin and other constituents During that These, with their high speed of 2,000 previous period of storage the applier can be kept either a misote, are capable of reducing fruit to a under cover or outdoors, provided that in the A mill of average size can deal with a ton of happ does not rest directly on the ground fruit in 10 minutes or less. Note of the never If that occurs, there is risk of an earthy taint presses installed are of the hydraulic type-being acquired and transmitted in due course. The milled fruit pulp—the "pomare"—is to the cider. Storage on a bed of stars or built up on the bed of the press into a "chees," rough boarding provides the necessary security as it is technically termed. The accessories used against this risk. From the foregoing it follows in juice extraction are devised to facilitate the that it is undesirable to mix in the same heap rapid and complete expression of the juice and of fruit apples of widely varying conditions of sits free drainage from the "cheese" of pulper repeness. Informatly of repness is essential if For this purpose a wooden slatted drainage. rack is first placed on the bed of the press When the fruit is adjudged to be fit for milling and on it is laid a netted or open-meshed cloth the practice in up-to-date factories is to wash of cotton or other suitable fibre. In this tho the phartie in dynamic and the force passing it on pomace is received and agreed evenly as far to the mill. This operation of washing is of as the edges of the drainage rick until a layer of relatively recent introduction into edge pomace about I am in this characteristic materials. The making routine and serves the double purpose overlapping ades of the cloth are then wrapped of the could be about the first and words of the country to the country to the country of the country to t of thoroughly cleansing the fruit and affording over the pomace layer and above this a second a convenient and simple opportunity for the irscl.splaced and the same procedure repeated remained d my apples which may have rotted until a "cheese" consisting of about severe during atorage. It has also been found that there or hayers of ponnee is obtained, the subsequent menagement of the fermenta-The press is then put in action and ton of the justice is easier and the clarification in the course of from 15 to 30 munites the belix tion processes rendered quicker and more of the available juice is expressed, the individual layers of pomace being compressed so tightly

in tubs fer from 12 to 24 hours before pressing. This maceration was found, when the older types of mills and presses were used, to facilitate the subsequent expression of the juice and was believed also to improve the vintage quality of the liquer. Occasional makers still adopt this practice but, with the modern machinery now widely used, it is doubtful if any advantago is derived from it. With the use of modern equipment it is possible to obtain from 75 to 80% of the original weight of the fruit as juice, if the fruit used is in good condition and at a suitable stage of ripeness. With older types of machines a yield of 70% must be ranked as above the average.

The residual pressed pomace after the expressien of the juice in the manner described contains about 70% of its weight in the form of juice which cannot he expressed hy a single pressing. Many makers break up the cakes of pressed pomace into a fine state of division hy a suitable machino and subject this pulverised pomace to a further pressing. By so doing an additional yield of about 20 gallons of juice per ton of fruit can be obtained. It depends upon the value of the particular juice in question whether or not this second pressing is economic.
With modern equipment and fruit in good

condition the average yield of juice per ton of fruit should approximate to 160 gallons.

In France, instead of the juice being expressed in this manner it is sometimes obtained by extraction from the thinly sliced fruit by a diffusion process similar to that used in the beet sugar industry. This method has also been tried by some English eider makers, but has not

been generally favoured.

The freshly expressed juice is a turbid liquid, somewhat viscous owing to its rather high pectin centent, and ranging in colour from a pale straw tint to a deep amber-brown. Its colour generally can be correlated approximately with its acid and tannin contents, acid tending towards the paler shades and tannin towards the deeper; thus a high-acid low-tannin juice is usually very pale and a low-acid high-tannin ene dark, while a high-acid high-tannin sample as a rule is decidedly darker than one of similar acidity but less tannin. Held in suspension in the juice are fragments of fruit tissue of varied size, the amount and character depending to some extent on the state of ripeness of the fruit at the time of milling and largely also on the degree of fineness of milling and the form of construction of the cheese and the kinds of cloths used. The coarser material is removed by straining the juice through a fine-meshed sieve after it leaves the press; but normally sufficient suspended solid matter remains in the juice after sieving to render it desirable to take other steps for its removal, so that the fermentation vessels may be kept in cleaner condition and risks of flavour depreciation due to its presence during fermentation reduced as far as possible. A fair clarification of the inice at this stage is mostly effected in practice by one or other of the following three ways. Centrifuging the juice usually is effective to a high degree,

rider makers prefer to allow the pomace to stand | factories possess appropriate centrifuges. Effective filtration of the juice would serve, but is difficult to attain, and also very slow, owing to the presence of pectins. A simple method is that of "keeving," which consists in allowing the juice to stand at rest for a period of several days (until fermentation hecomes active) in a large Defecation occurs, most of the open vat. suspended matter rising to the surface and forming a more or less dense head easily removable by skimming and the hulk of the remainder sinking to the hottom of the vat to form the lees, which are separable by racking at the appropriate time. The chief drawback to the "keeving" method is the acctic fermentation which invariably occurs, sometimes to a slight or even negligible extent when the juices are prompt in coming into active fermentation, hut at others to a degree leaving permanently in the prepared cider a distinct taint of aroma and flavour. To minimise this action keeving at low temperature (35°F.) has been adopted with some success where facilities for chilling are available. The most primitive method of juice clearing is still the simplest and the one to he advised usually when a centrifuge is not available. In this the freshly expressed liquor is placed at once in the fermentation casks and allowed to stand there with the bung open pending the onset of active fermentation. is associated in its earliest stages with the formation of a thick, sticky head of foam, which as formed carries with it much of the suspended material to the open bung-hole where it is extruded over the sides of the eask. Within a few days generally this phase passes, activo fermentation takes charge and then, for the first time, it is desirable to insert in the bunghole a hung fitted with a modern form of vent-tuhe which permits the evolved earbon dioxide to escape and prevents any atmospheric oxygen reaching the liquor.

After the preliminary clarification of the juice, by whatever method adopted, the fermentation process is allowed to proceed uninterrupted until it has reached the desired point. The great majority of cider makers allow the juice to fer-ment "naturally," in which case the wild yeasts which occur on the skin of the apple and are washed into the juice during the milling and pressing operations are the active microorganisms concerned. Other makers, however, control the nature of the fermentation hy the use of selected yeasts, cultures of which aro added to the freshly expressed juice in quantity sufficient to dominate the fermentation which place subsequently. More complete control can be obtained by the pasteurisation of the juice before the addition of the pure yeast culture; hut this method is open to the drawback that the pasteurisation of the juice affects

its flavour to some extent.

In the natural fermentation several different kinds of yeast are usually concerned. During the course of the production of the heverage three distinct phases in the micro-organism flora, characterised by the predominance of certain yeast types, occur. In the initial stages the juice usually is effective to a high degree, of fermentation years of the apiculatus type are but at present few makers outside the larger relatively numerous. During the most active

stages of fermentation which follow, yeasts of longer atorage period sometimes gives an the recersion, ellipsoideus, and Pastorianus types improvement in quality, but more usually are predominant. When active fermentation edicer is at 12 reak of quality during the late has been completed the residual flora is characterred by the presence of a relatively large proportion of small-relled veests of the Torula class. At this stage also there is generally an For draught eider e final filtration through a increase in the proportion of bacteria to seasts, unless the later stages in the production of the beverage are very carefully managed.

Two distinct rourses of procedure in connection with the fermentation stage are open to the eider maker. Under the older method the juice is permitted to continue to ferment until active fermentation romes to a natural eod. When that point is reached steps are taken to clarify by racking or other methods described below. With this method the ultimate character of the eider as regards sweetness depends only upon the extent to which the natural ferments tion of the juico is carried This varies greatly with different juires and is determined mainly by the content of soluble organic nitrogenous com-

pounds suitable for yeast nutration in the juice The alternative method open to the eider maker is the one of "checked fermentation," which has for its object the retention of a degree of netural sweetness in the liquor desired by the maker. This necessitates bringing active fermentation to a standatill when the corresponding point of apecific gravity is approached litacking for this purpose is unreliable, although occasionally stoppage of fermentation can be obtained in this way when the nitrogenous content of the juice is exceptionally low Filtration is far more effective, provided that pulp filters of a fairly modern type are used. The most recent way which has been found to give the maker a high degree of control-as well as other advan teges-is to centrifuge the fermented liquor when the desired point of specific gravity has been reached

In the production of sweet eiders the maker using the "checked fermentation" method obtains a beverage with a part of the natural sweetness of the apple retained, for which he must rely on the efficiency of his method of checking the fermentation Under the older method the degree of an eetness left in the liquor at the end of the fermentation is very variable striking differences being usually in sugar and in different cases and generally magnificant in falcohol content according to whether the product amount; the required ascetness has then to be provided by the addition of augir. For the production of dry ciders fermentation is allowed to proceed in any case until it comes to a natural end

At the end of the active fermentation period. vata used bitherto

opring and summer of its first season,

Before it is placed on the market for consumption some degree of refining is required. pulp filter is adequate; the same applies to bottled eiders intended for natural conditioning. Since, however, ciders refined in this way are subject to the risk of certain bacterial disorders. such as eider "sickness" and acetification, a baeteris-proof filter has been adopted in recent practice to obtain a sterile beverage, the liquor in that event has to be conditioned for

bottling purposes by artificial cerbonation, In bottling eider two methods are generally used The older method, in which the liquor is allowed to condition it elf naturally in the bottle by a slow after-fermentation, is somewhat uncertain in its results and has been largely displaced latterly by the method of artificial carbonation. In the former method cider reunires to be bettled at the correct time in it. life This varies according to the individual ciders concerned and considerable experience is required to determine when a given rider has reached the best time for bottling; if bottled too soon, the haud becomes avercharged with ges and an excessive deposit is formed in the bottle, if bottled too late, little or no ofter fermentation occurs in bottle, the hould accordingly remaining flat and being frequently disfigured by the development of 'likes' resulting from the growth of acctie liacteria. As a rule bettling under this method should be done between mid February and carly May in the spring following making. The carbonstion process of bottling has the great advantage that the cider can be bottled at any time of the year in condition for immediate consumption, while the problem of deposit does not ense, When ciders which have been sterilised by filtration through bacteria proof filters prior to bottling are used, all parts of the bottling plant as well as the bottles and cork must be steriled to prevent re infection with micro organisms

The chemical composition of mature eiders differs considerably in individual cases, the most

is of the sweet or dry type
The table on page 127 shows the composition of some typical ciders of various degrees of

aweetnres and dryness. Cider is subject to several ilisorders. Of these the commonest are arctification, rickness, whether this has been achieved artificially or ropiness and oiliness, and discoloration. The not, the liquor is placed in clean, tightly-bunged first three are directly iluo to the action of casks and allowed to mature for a period. For specific bacteria, and the last, in some instances the storage of order in bulk large reinforced at least, may be indirectly attributable to acidconcrete vessels lined internally with glass tiles destroying micro organisms. In the case of acetifior an impermeable, tasteless, and odourless cation, the "vinegar" fermentation, in which material unacted on by exter are being used to an several distinct kinds of acetic bacteria may be increasing extent and displacing the large wooden roncerned, all culers are attacked unless the liquor is protected et all stages of manufacture Generally the liquor reaches the period when and subsequently until ronsumed. Every sure it is sufficiently mature for consumption comes from the press more or less heavily during the course of the spring following the autumn or winter in which it was made. A exposure to air must be reduced to the lowest

1	A	В	С	D	E
Density	1008-5	1003-5	1000 0	1025 0	1019 0 %
Total dry extract at 100°	3 025	2 37	2 [°] 06	7-71	6-03
	1.67	0·23	0 -15	5-53	4-03
Alcohol	3·1	7-5	6·45	3 8	4·6
	0 050	0 05	0 036	0 073	0 074
Fixed acid	0 152	0 675	0 458	0 201	0 229
	0 041	0 170	0 058	0-208	0-234
A•h	0-175	0 375	0 215	0 248	0 235

during the active fermentation of the juice as possible should be kept saturated with this appearance. gar, by direct addition when necessary, throughout its life. Cider sickness, (the "maladic de la tourne" and "la pousse" of the French), is a violent form of alcoholie fermentation produced by the "sickness" hacillus and apt to attack sweet ciders, whether in eask or hottle, when the temperature rises during the late spring or thus—has been for many years a well known summer following manufacture. Its onset hy-product of the industry. In some countries appears without warning and is generally accompanied by an intensely rapid evolution of carbon dioxide. A characteristic odour and flavour are produced and in course of time usually a dense milky turbidity appears in the liquor in consequence of a eliemical action of formaldehyde and other aldehydes formed during the fermentation on the tannin of the eider. Ciders with a make acid content of 0 5% and over are not often attacked. Storage at the coolest temperature available is recom-mended for all sweet eiders which have not been passed through a bacteria-proof filter.

Ropines in eider is similar in character to the corresponding disorder of wines, and oiliness appears to be an incipient or little developed stage. It is not of frequent occurrence, but is specially apt to appear when sugar is added for sweetening purposes. The use of a hacteria-proof filter for the final elamineation of eider is probably the most effective method of prevention of this disorder. All sugar syrups used for sweetening should be sterilised before addition, as a precaution against infection from

Discoloration or "blackening" of eiders is an unsightly disorder, but otherwise not actually detrimental to the beverage. The affected hquor turns dark after exposure to air, in bad cases within a few ininutes of being poured into the glass for consumption. The discoloration is an exidation effect and associated with the presence of iron in a certain form in the liquor. The iron contamination is most often acquired from metallic utensils with which the pomace or liquor has been in contact, but

practicable minimum, for at no period does have had their acidity reduced by the action of cider attain a sufficient alcoholic strength acid-destroying hacteria appear to be highly sustorender it immune from the development of ceptible to the trouble. In some cases enzyme action also may be associated with its development. The use of sulphur dioxide as a preservative forms a natural protectant and the liquor as far gives some measure of protection against its

> During recent years several hy-products of the industry of some importance have been developed. In France a considerable proportion of the total output of eider each year is distilled for the production of potable alcohol. "Calvados"—a form of eider brandy obtained thus-has been for many years a well known eider vinegar (r. Vol. I, p. 46a) prepared by the controlled acetification of fully fermented eiders is produced extensively and is in considerable demand.

> Various uses for the pressed pomace left after the final extraction of the juice have been found. In its fresh state it has some value as a food for livestock, provided that it is not kept so long that acetification to a material amount occurs. The material is now generally dried in suitable drying machines until the moisture content is reduced to approximately 10%. In that form it is perfectly stable and can still be used for stock-feeding purposes, being frequently incorporated with other feeding stuffs of more concentrated food value in compound eattle cakes. Considerable quantities of commercial peetin are extracted from dried pomace at the present time and used in jam manufacture. B. T. P. B.

> CIGNOLIN. Dihydroxyanthranol, used in the treatment of various skin diseases (Bayer Products, London). B P.C

> CINCAINE. isoPropylhy drocupreme hydrochloride, used as a local anæsthetic Howards, London). B P.C.

> CINCHOMERONIC ACID. 3 4-dicarboxy he acid.

CINCHAMIDINE t. CINCHONA ALKALOIDS (this vol., p 162a).

CINCHENE r. CINCHONA ALKALOIDS (this vol. p. 15%)

CINCHOLOIPONIC ACID r. CINCHONA ALKALOIDS (this vol. p. 159d).

CINCHONA ALKALOIDS. DUCTION.—The first recorded instance of the certain soils are reputed to yield fruit specially isolation of a crystalline material from an susceptible to the disorder. A relatively high extract of cinchona bark is provided by the acidity in the cider tends to retard the appear"cinchonio" prepared by Gomes of Lisbon ance of blackening and those samples which in 1810. The basic character of this product initiated the investigations of these two French chemists, who showed that Gomes's preparation ronsisted of two alkaloids, enchonine and quinine (Ann. Chim. Phys., 1820, [11], 15, 292). Since then many other bases have been isolated from cinchona bark, the most recent being epiquinine and epiquinidine, isolated by Dirscherl and Thron (Annalen, 1935, 521, 48). The following list includes all the alkaloids 1850, gives a comprehensive acrount of the work

group:		the pari
Empirical Formula.	Name.	As a :
C, H,ON,	Paricine.	Java, I
C,H,ON,	Cinchonine, einchonieine (cin-	Were n
C19H24ON	Cinchotine (hydrocinchonine), cinchamidine (hydrocinchoni-	Java an
	dine), cinchonamine.	
C,,H,,O,N,	Cupreine.	country
C,H,O,N,	Ouinamine concilinamine	atrain (
C.H.O.N.	Quinine, epiquinine, quinicine	which it

quinidine. CaHAOANA Hydroquinine, hydroquinidine. C.H.O.N. Chairamine. conchairamine, chairainidine. conchatramidine.

C.,H,,O,N, Cusconine, concusconine, ancine Dieinchonine.

C₁₀H₄₄O₃N₄ C₄₀H₄₄O₃N₄ Composition Diconquiame, Javanine enscamine, cuscamiundetermined dine, cuscenidine.

It is deubtful whether these are all wellanthenticated aingle substances, as some of them are amerphous and not too well charse-terised. The alkaloids paytine and paytamine are sometimes included on the insufficient ground that " Pavta " bark (Aspidosperma sp.), from which they are obtained, has been used as a febrifuge,

Cinchona bark has a long and interesting history. It is a moot point whether its antimslanal action was knewn to South American Indians before the advent of Europeane; among travellers in that region, Arrot (Phil Trans. 1737-8, 40, 81) and others are of opinion that it was, while Humboldt and Markham held that it was not. The first cure of a European was reported by Jussieu and is supposed to have taken place in 1600, About 1638 the Countess of Chinchon, wife of the Spanish Viceroy of Peru, was cured of malaria by the use of the bark, and this event was rommemorsted by Linnwus in the name Conchona, which he adopted for the genus of plants yielding the drug.

The cinchonas occur on the western side of S. America between 10°N, lat. and 22°S. lat., i.e. in parts of Venezuela, Colombia, Ecuador, Peru and Bolivia. Though cinchona bark did not meet with universal acceptance among the medical profession on its introduction advent of a stematic planting, the rommercial into Europe about 1632, its use spread repidly varieties have been virtually redured to "factory and the great demand led to ruthless explortation bark" derived from Cinchona Ledgeriana or

was noted by Honton-Labillardiere, who told serious prospects of a cinchona famine. By Pelletier and Caventon about it, and thus that time the cinchonas had been classified botanically and it was feasible to rousider the formation of plantations in other parts of the world. A Dutch expedition under J. C. Hasskarl visited the einchona regions in 1852 to rollect plants and seed of as many cinchons species as possible, and in 1859 similar British expeditions were organised by Sir Clements Markham, whose book, "Peruvisa Bark," published in which can be regarded as belonging to this of these expeditions and of the early history of group;

result of these Dutch and British enternnchena plantatione were established in ndia, Ceylen and elsewhere, and at first nost successful in India, but in 1855 Ledger obtained a supply of seed of sa Calisaya, half of which was planted in nd half in India. This species did well in but failed in Indis, and in the former proved to be the source of a remarkable of trees, named Cinchona Ledgeriana. mproved by unremitting investigation on (quinotoxine), quindine, epi the part of the Dutch planters inte the effect of cultural methods on yield of alkaloid, gradually onsted India from the dominant position it seemed likely to take about 1880 and conferred upon Java the virtual monopoly in cinchons production which it enjoys to-day.

There are probably few drugs whose early history has been se fully recorded as that of canchons, and the reader interested in this part of the subject may be referred to the following sources of information additional to Markham's book already mentioned:

1860. "Nuova Quinolegia ef Pavon," Jehn Elet Howard.

1879. " Pharmacographia," Fluckiger and Hanpara. 1691. " Pharmacographia Indica 1891," Vol. 11.

Dymeck, Warden and Hooper, 1903. "The Commercial Producte of India."

Watt. 1908. "Scheikundige Budragen tet de Kennis der Java-kina, 1872-1907," van

Gorkem. 1922. Indian Trade Enquiry, Imperial Institute. Report en Cinchona Bark."

"Notes en the Cinchona Industry." 1931. Lecture to the Institute of Chemistry. B. Howard,

1931. " Proceedings of the Celebration of the Three Hundredth Anniversary of the First Recognised Use of Cinchons," Missouri Potanical Garden, St. Louis, U.S.A.

1932. " Kew Bulletin of Miscellaneous Information, 1932," No. 1.

So leng as the world was supplied with cinchons bark from the natural forests of South America a number of varieties of bark, distinguished by such names as "crown," red,"
"3ellow," "grey," "soft pitayo," "hard
pitayo," etc, were marketed, but with the

of the natural forests and by 1850 there were hybrids and "red bark" or "druggists' hark"

used for the manufacture of quinine and other einchona alkaleids, and the second for the production of tinetures, extracts, and other galenical preparations. Various devices have been introduced into the plantations to enhance the yield of bark, to raise its alkaleidal content, and to facilitate harvesting and preparation for expert, and information on these points will be found in a paper by D. Howard (J.S.C.I. 1906, 25, 97). Though the bulk of the world's supply of cinchona bark new comes from Java, small quantities are still obtained from S. America, India, Ceylen and San Thome, but experts from Jamaica and ether British possessions seem to have eeased. There are small plantacolony was German East Africa, from which included in Table I.

obtained from Cinchena succirubra, the first being small quantities have from time to time used for the manufacture of quinine and other been experted (Wersley, Bull. Imp. Inst. 1935, 33, 14).

It is of interest to compare the yield and nature of the alkaleids obtained from the old S. American barks with those of modern plantation bark, and that is pessible ewing to the existence of a few analyses made by modern methods on authentic specimens of old barks, which have been kept as museum specimens. These are included in Table I. For commercial and industrial purposes it is sufficient to determine the meisture, tetal alkaleids, and quinine content of cinchena bark. For that reason modern analyses recording the prepertiens of the feur chief alkaloids for a particular einchena bark tiens in Tanganyika Territory, formed when this are rare, but such as have been found are also

TABLE I.

Number.	Date of supply of bark,	Variety of bark,	e Total	59 Quinine.	5₹ Quinidine.	3. Cinchonidine.	% Cinchonine	e Amorphous et alkaloids.
1 2	1870 1870	C. officinalis (erown bark) C. pitayensis (hard pitayo)	5·48 6·66	3·09 2·42	0·14 0·82	1·34 0·19	0·32 0·98	0.59a 2.25a
3	1870 1865	C. Calisaya (yellow bark) C. Calisaya (yellow bark)	4·12 5·80	2·00 3·61	0·20 0·16	0.46	0.56	0.90a 0.93a
4 5	1914	C. Ledgeriana	9.10	7.92	0.08	0.10	0.08	0.93
	1919-1923	o. Deager tone	6.13	4.14	0.17	0.54	0.38	0.886
	(average)	"						
7	1925	,,	5.79	4.14	0.44	0.36	0.25	0.60c
8	1925	Hybrid bark	4.54	2.87	0.34	0.33	0.46	0.54c
9	1888	C. succirubra (red bark)	6.25	1.40	1	2.25	1.92	0.68d
10	1921	,,	6.07	1.14		1.43	1.73	1.76c
11	1930	,,	6.29	2.05	0.11	1.47	1.41	1.25f
12	1930	C. robusta	5.86	2.01		2.42	0.12	1.31f

(a) These analyses are from records of John Eliet Howard and the results must be accepted with reserve, as at the dates mentioned tho methods of separation of the different alkaloids

were very rudimentary.
(b) Heward, Trans. Roy. Sec. Trep. Med. Hyg. 1924-25, 18, 358. Average composition of Java bark, 1919-23.

(c) Gage, ibid., p. 350. (d) Hooper, "Year-book of Pharmaey," 1888, p. 430.

(c) Howard and Chick, ibid. 1923, p. 637.

(f) Barks from Tanganyika examined by Worsley, Bull. Imp. Inst. 1935, 33, 22.

superiority of Calisaya bark, even in 1865, as a in France and England, coupled with the source of quinine, and analyses Nos. 5 and 6 indicate clearly that Dutch caro of its lineal descendant, C. Ledgeriana, has improved the yield of alkaloids, the proportion of quinine present, and, what is equally important, has effected a reduction in the proportion of ein-chonidine, the alkaloid which persistently accompanies quinine, and whose control is the principal objective of most pharmacopæial tests for impurities in quinine sulphate.

In the early days of einchena planting, attempts were made, especially by de Vrij in

Comparison of these analyses establishes the followed by the manufacture of this alkaleid persistent advocacy by Meens in Java of cinchona cultivation for quinine production, soon secured for quinine a dominant position as compared with the other einehena alkaloids, which it still occupies and is likely to retain. Analyses Nos. 7 and 8 relate to "ledger" and hybrid barks produced in India, where these barks are still grown to meet the needs of the Government quinine factories in Madras and Bengal, which, however, also use considerable quantities of Java bark. The two analyses Nos. 9 and 10 serve to illustrate the striking Java, to popularise the total alkaleids of the difference in the relativo preportions of the lark as an anti-malarial drug, but the rapidity various cinehona alkaleids in C. succirubra with which the discovery of quinino was bark as compared with ledger and hybrid barks.

No. 9 is described by Hooper as that of a ide (sp.gr. 1-375). To this mixture, diluted if typical red bark, being the average of analyses necessary by adding a little water, 2 litres of of a large number of Indian grown samples petroleum are added, heated to 100° for 20 No. 10 is en analysis of an average red bark of minutes, with constant agitation; the oil is then Indian origin Nos. 11 and 12 are barks from decanted, and the extraction repeated with a Tangaoyika Territory. Partial analyses of second 2 litres The 4 litres of extract are "ledger," hybrid, and succirubra barks from Tanganyika Territory have been made by the Impenal Institute (Bull. Imp. Inst. 1918, 18, 337; 1920, 18, 22), by Greenish and Corfield ("Year-book of Pharmacy," 1923, p 652), and by Goodson and Henry (Pharm J 1930, 124, 351), and complete analyses by Worsley (c.), who also gives an account of the progress and

working of these plantations since 1918
As a matter of historical interest, it may be recorded that in the late nineteenth century a considerable quantity of cuprea hark was sold in the London market. This bark was obtained from Remijia pedunculata, a near relative of the cinchonas, and was remarkable in containing quimine and the corresponding phenol cupreme, some cinchonine and quinidine were also present.

but no emchonidine 2. EXTRACTION OF ALEALOIDS -The descriptions of processes quoted below refer both to extraction of the alkaloids for the purposes of analysis and as a first step in manufacture The underlying principles are the same for both objectives. The alkaloids occur in the bark in combination with acids, and it is necessary to liberate them from these natural sales in order to feethtato extraction by immuscible solvents such as ether, ehloroform, benzene, or petroleum The first step, therefore, is to grand the bara finely, usually to a powder that will pass a sievo is then recutened and mixed with an alkali, usually lime or soda time, and left to stand until the action is complete, after which the mixtura is dried, reground if necessary, and extracted with the selected solvent

separates, when it is run off and shaken with dilute sulphume acid, this acid solution is boiled, and while boiling is neutralised, to litmus crystallises out on cooling, while einchonidine, cinchonine, and quinidine remain in solution as aulphates. The quinine aulphate is purified by aiderably the more soluble.

Landrin (Compt. rend, 1889, 108, 750) gives weighed. the following description of a small-scale extrac-tion: 300 of bask are powdered and passed through a No. do arer, then mused thoroughly with little of mild. South time containing 15g, of quickline and 15g of solution of sodium hydron seed and 15 e.e. of 3% hydrochiora cell are paickline and 15g of solution of sodium hydron.

washed twice with 75 c.c. of 10% sulphuric acid added to 150 cc. of water; and a third time with one third of these quantities.

A recent example of a small scale extraction of "ledger" and hybrid bark in Tanganyika

Territory is quoted by Worsley (I.c.).
Schwyzer ("Dio Fabriketton der Alkaloide,"
Springer, Berlin, 1927) provides a detailed
account of a small scale extraction and a full account of factory operations with diagrams of plant (see also Vetter, Festschrift E C. Barell Basel, 1936, 541).

The following general processes may be quoted as typical of extraction methods for analytical purposes. The reagents mentioned are those prescribed in the phermacopouss, which must be consulted for particulars as to strength, etc. The "British Pharmacopera," 1932, proceeds as follows: 10 g, of bark in about No 60 powder are thoroughly mixed with 75 c.c. of strong solution of lead sub-acetate diluted with 125 c.c. of water, and set aside for 1 hour, 50 ce o ammoniscal alcohol are then added, the whole again mixed and left standing for I hour. The preparation is next transferred to a continuous extraction apparatus, where, with a little more of the ammoniacal alcohol, it is extracted for about 4 honrs The bulk of the alcohol is removed from the extract, 10c c of N sploburg acid and 40c c. with 40 meshes to the linear inch. This powder of water added, the mixtura heated to boiling, then allowed to cool and filtered through a tight plug of cotton wool, previously wetted, into a separating funnel. The residue in the flask is treated with 15 to 20 c.c. of boiling N/10. sulphume acid, and the acid extract, after cooling. Zebel (Chem Zig 1891, 15, 735) describes the also filtered into the separator. The flask and commercial process thus. The finely powdered the cotton wool plug are finally washed with cold bark is ground to a thin paste with lime, sodium water, acudified with dilute sulphurin acid, init hydrovide or sodium carbonate, and extracted alkaloids are completely extracted. 20 cc. of with ware paratim oil. On standing the oil chloroform are now added to the separator, the contents shaken thoroughly, and the chloreform run off into a second separator and any alkaloids removed from it by agitation with 5 cc. of N. paper, with sodium carbonate. Quinine sulphate sulphuric acid diluted with 15 c.c. of water, the chloroform being then rejected and the acid extract used to treat in the same fashion two further extractions, each with 20 e c. of ehlororecrystallisation from water after treating with form, of the contents of the first separator. This animal charcoal. The mother-liquor containing final acid extract is transferred to the first the other alkaloids is treated with sodium separator, excess of dilute solution of ammonia hydroxide and the precipitate of quinidine, added, and the alkaloid completely extracted emchaniding, and emchanine extracted with with aucressive quantities of chloroform. The dilute alcohol, which dissolves the first two, combined chloroform extracts are washed with a leaving cinchonino behind; the two former little water, the solvent distilled off, and the last can then be separated by means of their traces of chloroform removed by dissolving the neutral tartretes, that of quinidine being con- residue in 5 e.e. of 95% alrohol, which is then evaporated, and the residue dried at 100° and

honr. To the cooled mixture 200 e.e. of etherchloroform solution (chloroform 1, ether 3) and 10 c.c. of the stronger ammonia test solution are added; the well-stoppered flask is shaken mechanically for I hour, left standing overnight, and again shaken for half an hour next morning. The drug is then allowed to settle, 160 c.c. of the clear ether-chloroform liquor, representing 4 g. of drug, transferred to a separator, and the alkaloids completely extracted with 5% sulphuric acid. The acid solution is made strongly alkaline with ammonia test solution and the liberated alkaloids extracted with chloroform. The solvent is distilled off in a tared flask and the residuo dried at 100° and weighed. The weight multiplied by 25 gives the percentage of alkaloids in the bark.

The "British Pharmacopæia," 1932, specifics for the manufacture of galenical preparations the bark of C. Calisaya Weddell, C. Ledgeriana Moena, C. officinalis Linn., C. succirubra Pavon, and hybrids of either of the last two with either of the first two. The bark must contain not less than 6% of alkaloids, of which not less than half must consist of quinine and cinchonidine. The total alkaloids are estimated by the "British Pharmacopæia," 1932, process described above. In the total alkaloids so obtained "quinine and cinchonidine" are estimated together by precipitation as the mixed tartrates, from which the bases are recovered and weighed as such. Two examples of the tartrate precipitation process are given below and it is, therefore, nunccessary to describe in detail the "British l'inrinacopreia" method as used for the total alkaloids of the bark.

The "United States Pharmacopæia" (11th Rev., 1936) prescribes the dried bark of the stem or root of C. succirubra or its hybrids, or of C. Ledgeriana and C. Calisaya or hybrids of these with other Cinchona spp. The bark must contain at least 5% of alkaloids as estimated by the "United States Pharmacopæia" process described above.

The "German Pharmacopæia," 4th ed., 1926, specifies the bark of C. succirubra, and requires it to contain 6.5% of alkaloids calculated as quinine and cinchonine (mean mol. wt. 309 2) and determined by titration of the extracted total alkaloids with methyl red as indicator.

3. Analytical. (a).—Estimation of Individual Alkaloids. The pharmacopæial methods described above are not only examples of processes for the extraction of einchona alkaloids, they also provide modes of estimating the total alkaloids in a sample of bark. In the total alkaloids so extracted it is nearly always necessary to estimate quinine, or evinine and einchoulding, and sometimes each of the four chief alkaloids. It must be remembered that the estimation of one component in a mixture of organic substances can rarely be effected with the accuracy characteristic of the majority of determinations of constituents of inorganic mixtures. Of the numerous methods that have i been devised for dealing with mixtures of ein-

Organie Analysis," 5tb ed., vol. VII, p. 426). This method has the advantage that it provides an estimate of the possible yield of quinino sulphate from a bark, but it should be realised that when this method is applied to the total alkaloids of two barks, such as No. 5 and No. 9 in Table I, the quinine sulphate from No. 5 will contain less einchonidine sulphate than that from No. 9. Goodson and Henry (Pharm. J. 1930, 124, 351; Quart. J. Pharm. 1930, 3, 238; 1932. 5. 161) have examined the "quinine sulphate," " einchonidine tartrate," " quinidino hydriodide," and "einehonine" obtained in the application of this process to certain mixtures of einchona alkaloids, differing materially from those characteristic of the total alkaloids of "factory" bark and have shown that in such eases the precipitates obtained need examination. Howard and Chiek also point out in the description given below that the quinine sulphate obtained is not to be regarded as pure. The method does, however, give good results in skilled and experienced hands, and the in-experienced, before attempting it, should practise the process on einchona alkaloid mixtures of known composition.

To obtain satisfactory results by Howard and Chick's method a quantity of alkaloid of about 5 g. should be available. It is desirable to start the assay with the estimation of the total alkaloids, as this simplifies the subsequent work and the result is a useful check on the estimation of the individual alkaloids. For this purpose it is only necessary to ovaporate to constant weight the extract obtained by any of the methods described under the previous heading; or, in the ease of the paraffin and sulphuric acid extraction, to make alkalino with sodium hydroxide, wash ont with chloroform, evaporate to dryness, and dry to constant weight at 100°-110°. Having obtained the weight of total alkaloids, add 3 c.c. of 10% sulphuric acid for every gram of alkaloid, and warm on the steam-bath till completely dissolved. Then add 80 e.e. of distilled water for every gram of alkaloid, boil, and while boiling neutralise exactly to litmus paper with dilute ammonia solution (10%). On cooling for 2 or 3 hours with occasional stirring practically the whole of the quinine will crystallise out as sulphate, owing to its being almost insoluble in ammonium sulphate solution. Filter through counterpoised filter papers on the filter-pump, press the crystals down well, pumping as dry as possible, and following with two washes of cold water each about equal in volume to the bulk of the precipitate. The filter papers and erystals are now dried to constant weight at 100°. The weight of anhydrons quinine sulphate found multiplied by 1.18 gives the amount of rough hydrated quinine sulphate, or by 0.868 the amount of rough quinine alkaloid. This quinine sulphate is neither white nor pure, being usually contaminated with colouring matter and einehonidine sulphate, and requires further treatment. But with the present-day Ledgeriana barks the weight may be taken as representing the true amount of quinine sulphate. chona alkaloids, probably the best is that as in some thousands of analyses of this bark of Howard and Chick with a slight modification by B. F. Howard and O. Chick, the amount intreduced by Chick (Allen's "Commercial of quinine sulphate left in the mother liquor

almost exactly compensated for the impurities in | this definition to "totaquina is a mixture of

the crystals weighed. To the mother liquor and washings from the

solved in 20 c. o. of hot water, and and reconceased by less than 70%, of crystallisable enhanced during 2 or 3 hours. Canchondene is completely prespirated as attracts. This is collected and the contemporary of the contempora the bark contain very little emchonidine, and no tartrate be obtained after standing 3 hours.

to ensure complete precipitation. The filtrate is made alkaline with sodium hydroxide solution and shaken out three times with other, using 100 c c of washed other each time. The combined ethereal extracts are in turn extracted with three lots of 10% sulphure acid using 10, 10, and 5 e c. respectively, and the acid liquids transferred to a separator containing 25 e.c. of ether Sufficient sodium hydroxide solution is then added to render the liquid distinctly alkaline and the whole well shaken and left to stand I hour. Conchemne crystallises out and is collected by filtration. The aqueous layer is shaken out twice more using 25 c c of ether each time. The crystalline conchoning is washed with the second lot of other, then dried and weighed, and 0 08 g added to correct for the solubility of the alkaloid in ether. The combined ethereal liquids are extracted with acetic seid, the extract neutralseed at the boiling-point with 10% ammonia solution, using sensitive litmus paper as an indicator, 5 g of solid potessium sodied added, and the liquid allowed to cool and left 2 to 3 hours to enable the quinidine hydriodide to 40 c.c. of solution of sodium hydroxide and 80 c c. separate. The clear supernatant higher is of chloroform and set aside, with occasional deranted through counterpoised filter papers, the precipitate is warmed with a little 50% alcohol to dissolve amorphous alkaloidal hydrodides, the whole being then filtered and the precipitate washed with small measured amounts of 50% alcohol until the quinidine are washed with a little water, the selvent hydrodide is pale yellow in colour. The precipitate is dried, weighed, and 0 0008 g added to the weight for each c e. of 50% alcohol used. The corrected weight multiplied by 0 717 gives the amount of quinidine alkaloid present. The filtrate is evaporated nearly to dryness with the addition of ecough dilute sulphurie acid to keep the remaining bases in solution, transferred to a separator, made alkabne, and the amorphous bases extracted three times with other using 25 e e. each time. The othereal stantially that of Howard and Chick described extracts are evaporated to dryness and the above, and the anm of the percentages of these readue weighed as amorphous alkaloids.

Totaquina -Estimation of the components in mixtures of cinchona alkaloids has assumed

alkaloids from the bark of Cinchong succirulera Pavon, Canchona robusta Howard, and other suntable apecies of Canchona. It contains not the required standard (Type B). These two types of totaquina differ considerably in comit is best to leave the solution for a full 24 hours position, but the pharmacopicial process is designed to deal with both.

"British Pharmacopana" Method for Totaquina -A mixture of 2 g of totaquina with 20 e c. of N-sulphuric acid, 40 e c. of water. and 40 ee of 95% alcohol is heated to boiling and, while still bot, made just alkaline to litmus by addition of N/10 sedium hydroxide, To the cooled liquid N/10 sulphume acid is added drop by drop intil the solution is slightly and to litmus the liquid is then boiled for I or 2 minutes, cooled, and rendered slightly acid to litmus if necessary, boiled, and filtered into a tared flash, the original vessel and filter being washed with boiling water until all the alkaloid is extracted. The washings are added to the filtrate, which is then evaporated to about 120 g . 30 g of powdered Rochelle salt are added, the mixture shaken to complete solution and set aside for 24 hours. The precipitated tartrates are collected on a haedened filter, the flask and filter being washed with 80 c c. of a 20% and and there being wanted with ab c c. of a 20% (weight/volume) solution of Rochelle salt used in suitable portions. The filter and washed precipitate are returned to the flask, shaken with agitation, until complete solution is effected. The chloroform solution is then separated, the ilask and the aqueous hound being washed with ehloroform in portions until all the alkaloids are extracted. The combined chloroform solutions evaporated, the last traces being removed by dissolving the residue in 5 e e, of alcohol, which is then evaporated, the final readus being dried and weighed as quinine and cinchonidine. In this residue the amount of quinine is estimated by a methoxyl determination using 02 g; 1% of methoxyl is taken as equivalent to 10-45 of quinine. The procedure for the estimation of canchonine and quandine in the filtrate from the quinine and cinchonidine tartrates is subfour alkaleids found is taken as the percentage

of crystallisable alkaloids. Instead of estimating quinine and einchonidine greater importance owing to the introduction by by either of the two processes described above, the Malsina Commission of the League of Nations the two alkaloids may be precipitated together of "totaguno," defined as containing at least as tartrates and the content of each deter-10% of crystalline askalonds of which not less mined polarimetrically. This plan forms part of than 10% must be quinne, with not more than the following process devised by Howard as 20% of amorphous alkaloid, and mosture and particularly suitable for einhone fabrings (see mineral matter not more than 5% each. The Quinetum, and also for totaquinas of type I. British Pharmacopcus, "1032, has modified and was adopted as an alterostic method by the Commission of the League of Nations to investigate the standardisation of totaquina (CH)

Malaria 183, Geneva, 10/10/1932).

Committee Method for Totaquina .- Ten grams of the product are dissolved in 50 c.c. of 10% hydrochloric acid, the solution being transferred to a large separator containing 750 c.c. of washed other. Excess of ammonia solution (sp.gr. 0.059) is added and the mixture shaken vigorously. The crude cinchonine which remains undisselved is filtered out at the pump, the filtrate separating into two layers, (a) ethereal, which is returned to the first separator, and (b) aqueous, which is placed in a second separator, where it is re-extracted with another 500 c.e. of ether, which is added to the first lot, the aqueous liquid being reserved for further treatment. The erude einebonine is washed, first with 50% alcohol and finally with acctone, until the washings in each case come through nearly colourless, after which it is dried and weighed. The washings are evaporated to dryness and the residue of amorphous alkaloids weighed. The combined ethereal solutions are washed out three times with 5% tartaric acid solution using 100 c.c., 50 c.c., and 20 c.c. conscentively; the united acid liquids are heated to boiling, neutralised to litmus paper with 20% sodium hydroxide solution, set aside for 18 hours to allow completo precipitation of quinine and einehonidine tartrates, and the precipitate collected, washed with as little cold water as possible, dried, and weighted. In it quinine and einchonidine are determined polarimetrically by the method described below.

The filtrate is transferred to a separator, excess of 10% ammonia solution added, and the mixture extracted twice with other using 500 e.e. each time. The aqueous liquid is added to the original aqueous liquor already reserved. The combined ethereal solutions are extracted three times with 2.5% sulphurie acid using 80, 40, and 30 e.e. conscentively; these acid extracts are mixed, rendered alkaline with 10% ammonia solution, shaken vigorously with 50 e.e. of washed ether, and the mixture left standing 24 hours to allow a further quantity of cinchonine to crystallise out, which is collected, dried, weighed, and the weight added to that of the cinchonine already isolated. The filtrate is shaken out with two further lots of ether. 500 c.c. each, and the aqueous liquid added to the two aqueous liquors already reserved. united ethereal extracts are then shaken out three times with 2.5% sulphuric acid using 50, 40, and 30 e.e. consecutively; the mixed acid extracts are heated to hoiling, neutralised to litmus with 20% sodium hydroxide solution, excess of powdered potassium iodide, about 10 g., added, and the solution left standing overnight. The liquid is then poured off and added to the reserved aqueous liquids; the precipitate is well stirred with 20 e.e. of 50% alcohol and the mixture warmed to about 50°C, to dissolve amorphous hydriodides. The quinidino

Committee of Experts appointed by the Malaria | weight multiplied by 0.717 gives the amount of quinidine base.

The bulk of the alcohol is evaporated from the filtrate, the residue of the latter added to the alkalino liquers already reserved, and the whole extracted with a mixture of benzene and amyl alcohol (3:1) ("British Pharmacopæia," 1914). This extract is then evaporated to dryness and the weight of the amorphous alkaloids left added to that found in the initial extraction

to give the total amorphous alkaloids.

Polarimetry of Tartrates.—If the precipitate of tartrates of quinine and einchonidine is too dark-coloured to be used as such for polarimetric reading, it is recrystallised by the following process: About 0.8 g. of the precipitate, dried at 90° to 100°, is mixed with 15 c.c. of a cold, saturated, aqueous solution of quinine and cinchonidine tartrates, 1.5 g. of tartarie acid added, and solution effected by boiling, 0.02 g. of decolourising earbon being added. The solution is then filtered, about 5 e.e. of a cold, saturated, aqueous solution of quinine and cinchonidine tartrates being used to wash the earbon on the filter. The filtrate is raised to boiling point and at that temperature neutralised to litinus by sodium hydroxide selution and set aside with occasional stirring. The purified tartrato precipitate is filtered off, washed with 3 lots each of 5 e.e. of cold, saturated, aqueous solution of quinine and einchonidine tartrates, then with 5 e.c. of distilled water and finally dried at 90° to 100°C, and weighed.

For the polarimetrie determination 0.5 g. of tho purified tartrate is dissolved in 5 c.c. of Nhydrochloric acid, made up to 20 e.e. at 17°C. with distilled water and the optical rotation read in a 2 dem. tube. Pure quinine tartrate under these conditions gives a reading of -10° 50' and each decrease in this reading of 2° 65' represents 1% of cinchonidine tartrate. The weight of tartrate × 0.80 = weight of free alkaloid.

(b) Examination of Commercial Quinine Salts.-Most pharmacopoins specify standards for quinine salts and prescribo tests with which the salts must conform. The principal impurities to be dealt with are einchoniding and hydroquininc. The latter is not of practical importance as it is now well established that hydroquinine is at least as effective in the treatment of malaria as quinine, and as it is more costly it is not likely to occur in quinine in abnormal quantities. Much labour has been expended in devising methods for the detection of other alkaloids, and particularly einchonidine, in quinino sulphate.

Kerner's Test.-Many of these tests are of little more than historical interest and most pharmacopæias now rely upon either Kerner's test or a polarimetric test to ensure the absence of an undue proportion of einehonidine. Kerner's test (Z. anal. Chem. 1862, 1, 150; Areb. Pharm. 1878, 14, 438; 1887, 25, 712, 749) depends upon the fact that quinine sulphate is the least soluble of the sulphates of the cinchona bases; and that quinine is the most soluble of the common hydriodide is then filtered off, washed as einehona bases in dilute ammonia. When 2 g. economically as possible with warm 50% of pure quinine sulphate is shaken with 20 c.c. alcohol until apparently free from amorphous of water at 60° for some time, and then cooled hydricolides, then dried and weighed. The to 15° for at least half an hour, to prevent super-

saturation, it gives a filtrate which precipitates with occasional shaking. Then filter through a with aqueous ammonia (1997, 9539), the per-filter-paper of 8 to 10 cm, diameter and transfer cipitate just redissolving when 43 e. of dibute 5 e. of the clear filtrate at 15° to a test tube ammonia solution have been added. Quinne and mix at gently (without shaking) with 6 c. culphate containing any of the other cinchona of ammonia test solution containing not less sulphates, when treated in this manner, requires than 10% and not more than 10 2% of ammonia; more ammonia to give a clear solution; and by the ammonia solution must also have a temperathis means some idea of the purity of the ture of 15° and be added all at once, a clear quinine sulphate is obtained. The details of liquid should be produced the test as prescribed in different pharmacopcers vary considerably, and there has been much discussion as to the best form Quinine sulphate containing about 5% of cinclionidine and hydroquinine aulphates will require between 6 and 7 cc of ammonia, while one containing about 1000 of these impurities will require about 10 c c. Points to be noticed in applying the test are that the weights of oninine sulphote and water taken do not, within reasonable limits, affect the result , the solution should be cooled gradually to 15°, but not below, and retained at that temperature for not less than 30 minutes, the filtration should be rapid and the rowing carefully done by gently rotating and tilting the testing tube, otherwise clotting of the precipitate may take place and a false result be obtained. This test is greatly influenced by the presence of free quinioe, when the result may be much in excess of that required for a neutral sulphate of the same degree of purity (D Howard, Pharm J 1896, [1v], 3. 505), and also by the presence of alkaline sulphates, which make it appear purer than it is (Tutin, Pharm J 1909, [iv], 29, 606) These points should be ascertamed by testing the filtrate for neutrality with hitmus paper, and by evaporating 5 e c to dryness and constant weight when the residue should not weigh more than 0 00% g., showing the absence of soluble inorganic salts (Biginelli, Monit Seient 1908, 22, 175) Should the cooling of the mixture be carried out above 15° a larger amount of ammonia will be required, and between 15° and 20° a correction of 0 5 e e of ammonia for each

degree above 15° may be applied. The following is the form of this test adopted in the "British Pharmacopens," 1933 1 g of quinina aulphate, which has been dried at 50° for 2 hours, is boiled with 30 ce of water for l or 2 minutes under reflux in a 100 e c resistance glass flask. The flask is cooled to 15° by means of water at 15°, then corked, shaken vigorously, and Lept at 15° for 30 minutes with frequent shaking. The contents are tiltered rapidly through a filter-paper 8 to 10 cm in diameter 5 e c of the clear filtrate are trans ferred to a test tube ; to this are added all at once 65 cc of a solution of ammonia, which must contain not less than 100 (w/w) and not more than 10 2% (w/w) of ammonia, and have a temperature of 15° The contents of the tube are then inixed gently without shaking.

A clear bound should be produced at 15° The 'United States Pharmacopoeia" (11th Rev. 1936) prescribes the following form: Shake 1 5 g of quinne sulphate, which must be neutral or faintly alkaline to himus paper and lave been dred at 50° for 2 hours, with 20 ce. of distilled water at 65° for 30 minutes, cool to 15" and set uside at this temperature for 2 hours i

Kerner's test is designed for use with ouining sulphate and in order to apply it to any other salt of quinine, that salt must be converted into neutral quinine aulphate without change in the composition of the base. Until recently the methods of achieving this conversion left much to be desired. The much improved method adopted in the "British Pharmacopera," 1932 may be summarised as follows, pung the case of quinine hydrochloride, B HCl.2H.O. as an example

Ommne hydroel loride (1 2 g) is dissolved in water 450 c c.1 containing dilute sulphuric and (5 ec) Dilute ammonia solution (5 ce) is added and the mixture extracted successively with 30 and 20 be of chloroform Each chloroform solution is slinken twice with water. ueing 10 c c. each time, and using the same water for both lots of chloroform The mixed chloroform solutions are st once evaporoted quickly on a water hath, most, but not all, of the chloro-form being removed. The list traces of chloroform are removed by dissolving the residue in 3 or 4 c c of a beolute alcohol and again evaporating quickly on a water bith, with the help of a current of air, until a white opaque residue is obtained. This is dissolved in 20 cc. of alcohol, 20 ce of water added, and I ce of a 0 02% (w/s) solution of methyl red in alcohol (90%). The colution is heated to 75° and at that temperature N/5 sulphuric acid is carefully added, until the colour matches that of a solution at 20° prepared by mixing 5ff c c of the "British Pharmacopæia" solution of pu 5 11 with I ce. of a 0.02 solution (w/v) of methyl red in alcohol (90%). In this neutralisation processionly acid must be used and any necessity for the addition of alkali must be avoided. The neutral liquid is then evaporated to dryness in a porcelain or resistance glass dish on a waterbath, and the dry residue powdered and used for the Kerner test as described above.

In the "United States Pharroacopæia" (11th Rev) the mode of application of the Kerner test to quinine base and salts, other than the sulphate, is varied Quining base is neutralised to sulphate by dissolving 1.5 g. of the base dried at 100° in alcohol (25 cc) and hot water (59 cc), adding enough X sulphune acid (about 5 cc) to make the solution acid. and then titrating back to neutrality with A section hydroxide, using methyl red as indicator. For the acid sulphate 252 g, of the salt B H₂SO₄.7H₂O is neutralised to methyl red with N section by drovide solution. With quinine dihydrothloride, the base is recovered from 18g of the salt B 2HCI, and converted to sulphate as described above for quinine base. For quimmo and urea hydrochloride quimme, base is also recovered from 3 g. of the substance

C,H,2O,N, HCI,CO(NH,), HCI,5H,O

as described above for quinine base. The neutral aqueous alcoholic solutions of quinine sulplinte with some sodium sulphate produced in the foregoing processes are evaporated to dryness and the powdered dry residue used for the Kerner test as described for quinine

sulphate.

quinine The pharmacopoial standards for salts vary considerably, as indicated by the different amounts of ammonia solution allowed for the production of a clear solution: 6.5 e.c. in the "British Pharmacopæia" (1932), 6.0 in the "United States Pharmacopæia" (XI, 1936), 4.0 c.c. in the "German Pharmacopæia" (VI, 1926), and 4.5 c.c. in the "Dutch Pharmacopæia" The 1926). last-mentioned conœia (V, pharmacopæia accepts the Kerner test for quinine sulphate, but the general method for quinine base and salts is a polarimetrio test, are given in the following table :

and 1.5 g. of the dried recovered base neutralised | based on the allowed optical rotation, -17.7° to -18.0° for a di-milligram-molecule of an hydrous quinine base in a calculated equivalent of a mixture of sulphuric and hydrochlorie acids, the solution being diluted to 20 e.e. with water and read in a 2 dem. tube. Thus, in the case of quinino sulphato 782 mg., i.e. a molecular weight in mg. of the salt

$$(C_{20}H_{24}O_2N_2)_2 \cdot H_2SO_4, 2H_2O$$

equivalent to 746 mg. of anhydrous quinine sulphate, (C₂₀H₂₄O₂N₂)₂·H₂SO₄, or to 648 mg. of anhydrous quinine base, is dissolved in water with the aid of 0.5 c.c. of 4N-sulphuric acid and 0.5 e.e. of 4N-hydrochloric acid and the solution made up to 20 c.c. and the optical rotation observed in a 2 dem. tube. The equivalents of salt and acid used in this and some other eases

Pharmacoporial substance and formula.	Equivalent wt. for 648 mg. of anhydrous base.	Acid to be added.	Optical retation in a 2 dem. tube.
Quinine base, C ₂₀ H ₂₁ O ₂ N ₂ . Quinine hydrochloride, C ₂₀ H ₂₁ O ₂ N ₂ ·HCl, 2H ₂ O. Quinine sulphate, (C ₂₀ H ₂₁ O ₂ N ₂) ₂ ·H ₂ SO ₄ , 8H ₂ O. Quinine acid sulphate, C ₂₀ H ₂₁ O ₂ N ₂ ·H ₂ SO ₄ , 7H ₂ O	0.7208 g. of anhydrous salt 0.782 g. of salt dried at	0.5 c.c. 0.5 c.c.	-17.7° to -18.0° at 20°. This value must be lowered by 0.023° for each degree of temperature above 20°.

for by dissolving the quinine sulphate in dilute sulphuric acid, cooling to 0° and adding a 4% aqueous solution of potassium permanganate so long as rapid decolorisation takes The solution is then warmed, manganese dioxide filtered out, washed, and the filtrate and washings made alkaline and the liberated hydroquinine extracted with ether, the solvent distilled off, and the residue dried to constant weight. Processes less open to obvious objection are catalytic hydrogenation in an apparatus arranged for quantitative work, such as that described by Paget and Solomon (J.C.S. 1932, 1198) and used by Buttle, Henry and Trevan for the estimation of hydro-bases in commercial cinchona alkaloids (Biochem. J. 1934, 28, 426) or the mercuration process of Thron and Dirscherl (Annalen, 1935, 515, 252), which depends upon the fact that each of the four primary cinchona alkaloids, in virtue of the presence of a vinyl group in the moleculo can combine with a molecule of mercuric acetate, while the hydro-bases in which the vinyl group has been reduced to an ethyl group cannot be mercurated in this fashion. The process is carried out as follows: 1 g. of the base (or an equivalent quantity of a salt) dissolved in 6 g. of 10% sulphuric heid is treated with 14 g. of a 10% solution of mercuric acctate in 5% acetic acid, and the mixture warmed for 4 hours at 40° to 50". When cold, 10 g. of ammonia

Hydroquinine has until recently been tested hydro-base extracted with other; usually 20, 10, and 10 c.c. in successive portions are sufficient. The combined other extracts aro washed with water, the solvent distilled off, and the residue of hydro-base dried at 105° and weighed. The alkaline mother-liquor is acidified with sulphuric acid, 0.5 g. of phosphorous acid added, and the solution heated to boiling for a short time. The clear liquor is then decanted from the mercury formed, the latter washed with water, and the mixed liquor and washings made alkaline with ammonia and the quinine extracted with ether.

(c) Examination of Other Commercial Cinchona Alkaloids.—Apart from quinine the only cinchona alkaloid of therapeutical importance is quinidine, of which the sulphate is included in the British and United States "Pharmacopæias." In addition to qualitative tests, c.g. the thalleioquin reaction, both pharmacopmias require that (a) 1 g. of the salt shall be soluble in 5 c.c. of a warm mixture of chloroform (2 vols.) with absolute alcohol (1 vol.), and (b) that when 0.5 g. of the salt in 15 c.c. of boiling distilled water is mixed with 0.5 g. of neutral potassium iodide in 5 e.c. of distilled water, the mixture being then cooled to 15° and kept at that temperature for 1 hour with frequent agitation, the filtrate on addition of 2 drops of dilute solution of ammonia does not become turbid within I minute.

The four primary einebona alkaloids are solution (sp.gr. 0.96) are added and the liberated I divisible into a layorotatory pair, quining and

emebonidme, and a dextro-rotatory pair, Marschall (Annalen, 1911, 382, 360). A spectro-quinidma and cinchomme. In commercial graphic method of determining cunchons alka-specimens the two components of each pair loids has been proposed by van Arkel and van usually contaminate each other, but not the der Wielen (Pharm. Weekbl. 1935, 72, 1198; components of the other pair. Each of the four 1937, 74, 1514). alkaloids is usually accompanied by its hydro base, and in cinchonine the amount of hydro-the usual alkaloidal reasents and precipitants lase emchanine may be as much as 140 and in Alkatoins). Characteristic of the methorylated quinidine up to 20%.

The author has not been abla to find in the literoture any satisfactory systematic method for the examination of commercial samples of quinidine, cinchonine and cinchonidine, but suggests that, bearing in mind the facts quoted above, it should be easy to evolve such a method when required, eq in (1) cinebonidine, the probable impurities are (2) quinine, (3) hydroguinine, and (4) hydroguinehousement. The vinvl bases (1) and (2) can be separated quantita tirely from the hydro bases (3) and (4) by Thron and Dirscherl's mercuration process (see above), and a methoxyl determination in (1)+(2), and (3)+(4) thus recovered will give the amount of quinine and hydroquinine present respectively, or the amount of each component in each mux ture may be arrived at by a polarimetric The author has found it conestimation. venient in dealing with mixtures of cinchona alkaloids to determine the rotation for a M/40 solution of the base in N/10 sulphuric acid The specific rotations for carefully purified specimens of the eight principal cinchona alkaloids under these conditions are as follows (Buttle, Henry, and Trevan, ! c.) :

The figures are for anhydrous base in each case, but the equivalent of air dry base is weighed out for the determination

4 GENERAL PROPERTIES OF THE CINCHONA ALKALOTOS -The eight principal cinchena alkalords alore mentioned (i.e. einchonine, quining, their stereoisomerides and dibydroderivatives) are white, odourless but intensely bitter, high melting, erystalline solids. bases are almost insoluble in water or aqueous solutions of salts and alkalis, but dissolve, in general, in dilute mineral acids and in a variety of organic solvents

and may even destroy, its florescence. For a direct smalled gives most of the emchonatories qualitative connection between the appearance of interescence and the chemical constitution of the alkaloids and their derivatives see Rales and

The enchona alkaloids respond to tests with alkaloids are the thellesoguin and crithroguin reactions. The former (Brondes, André) is carried out as follows: to an aqueous solution of, for example, a quimina salt, or to grimme dissolved in dilute acctic acid, is added a slight excess of chlorina or bromina water or a solution of bleaching powder, followed by an excess of ammonia giving risa to an emeroid green coloration or precipitate, depending on tha amount of quinina present. The reaction is said to be weakened or inhibited by caffeine, antipyrine and other substances. The erythroquin reaction (Vogel, Abensour) is carried out in a similar fashion, except that before adding the ammonia, a drop of 10% potassium ferroor ferrs eyansde is added, the coloration produced is red: this reaction is said to be even more delicate than the thalleroous test. The most delicate test however is that afforded by

the fluorescence already mentioned The cinchona slkaloids are strongly optically active. They fall into two groups of four, the dextro- and lavo rotatory series respectively. To the former belong cinchonine, quinidine, and their dibydro derivatives; einchonidine, quin ine, by drocinchonidine, and hydroquinine are invorotatory. In general, the rotations of the bases in alcohol are of the same order of magnitude as those of the neutral salts (BHA) in water. The rotations of the saits, however, progressively increase in arithmetic value with addition of acid. A maximum is reached at about the stage at which the amount of seid added auffices to convert the neutral salt into the acid salt (B 2HA); with further additions of acid the rotation alowly falls. This charac-Tersite behaviour was first observed by Hesso (Annslen, 1873, 165, 217. See also Oudraman, 1842 1872, 33; 1870, 1974, 45; 1881, 209, 33; Rec trav. chum. 1882, 1, 18; Liquier, Compt. rend. 1926, 183, 195; Detzel and Söllner, Arch. Pharm. 1930, 288, 629; Lapp. Compt. rend 1932, 195, 243; Bull. Soc chim. 1935[1], 2, 1407; Emde, Helv. Chim. Acta, 1932, 539, Ludwiczakowna and Suszko, Bull. Acad. Polonaise, A. 1934, 402).

The cinchona alkaloids are atrong discidio bases, which form two series of asits, viz. neutral The four alkaloids which contain a methors 1 saits B HA (called "basic" saits in Fronce), group (quinine, etc.), dissolved in much dilute and acid salts B 2HA (in France "neutral" sulphuric, nitric, or other oxy-acul, exhibit a salts) B 4HA is also occasionally met with. blue fluorescence, which is particularly intense. The neutral salts are neutral or faintly alkaline when viewed in ultra violet light and allows of to himus; the acid salts are acid to litmus but the detection of minute amounts of quinine (see, neutral to methyl orange or Congo red. These e g Canals and Peyrot, Compt rend 1934, 193, are, in general, colourless, crystalline solids, 746). Solutions in the halocen hadracids are which tend to become discoloured on exposure not fluorescent; the addition of such an acid to light. It has been claimed that exposure of to a solution of minime in an exy-acid weakens, einchona alkaloid salts in aqueous solution to

just mentioned, formed when the acid sulphates or tartrates are heated to melting, result also on se mild a treatment as boiling in dilute acctic acid solution, which, indeed, is more certain of its effect than the use of mineral acids. With more concentrated mineral acids the unsaturated alkaleids undergo complex isomerisation and transformation (see Section

5d, p. 149).

In temperate climates the cinchena alkaloids are used mainly as bitter tonics and anti-pyretics, but the great hulk of the quinino manufactured is used in tropical countries as a specific for the treatment of malaria. In common with the other cinchona alkaloids it exerts a direct toxic action chiefly on the asexual forms (schizonts) of the malarial parasite. These substances, therefore, provide an example of chemetherapeutic action, discovered and applied long before the principles upon which chemotherapeutic investigations as now conducted were laid down by Ehrlich. For over 300 years, at first in the form of cinchona bark or its galenical preparations, and in the last century as the alkaloids or their preximate derivatives, cinchena has been the sele remedy available against malaria.

As knowledge of the etiology of this diseaso increased it became evident that these alkaloids are not complete remedies and, in particular, it was realised that they were relatively ineffec-tive against the sexual forms (gametocytes) of the malarial parasite. Within the last decade it has been found that complementary drugs exerting gametocidal action can be synthesised, of which 8-(w-diethylamine-isoamyl)amine-6methoxyquinoline (plasmoquine) is the best knewn example. This substance resulted from the joint chemical and biological investigations of Schulomann, Schönhöfer and Wingler (Klin. Woch. 1932, 11, p. 381) with Rochl (Arch. Schiffs. Trop. Hyg. 1926, 30, Beiheft 3, p. 11; 1927, 31, Beiheft 1, p. 48). An interesting development of this work was schizzed by Misterland M. achieved by Mictzsch and Mauss (Klin. Woch. 1033, 12, 1276; Angew. Chem. 1934, 47, 633), who in conjunction with Kikuth (Deut. med. Woch. 1932, 58, 530) found that the same side-chain, inserted in position 5 in S-chlere-3-methoxyacridine, produced the drug new known as "atcbrine," which like quinine is schizonticidal in action. These successes have led to great activity in this type of investigation, not only in Germany but in this country, France, Russia and elsewhere. There are numerous other minor applications of the einchona alkaloids in medicine, e.g. the use of quinine-urethane as a selerosing agent, of quinine-urea as a local annethetic, and of quinidine in the treatment of auricular fibrillation. Quinine and certain of its derivatives, such as the Morgenroth compounds referred to later, are powerful bactericidal agents, and though the high expectations at first based on these results have not materialised in practice, hope has not yet been given up that it may be possible to develop practicable, pneumococcicidal agents with one or other of the cinchons alkaloids as a starting point and work on these lines is being intensively pursued, especially in Japan and the United States.

5. CONSTITUTION OF THE CINCHONA ALKALOIDS.—(a) Structure. The principal alkaloids of this group have been shewn to have the following structure:

in which R may be hydrogen or methoxyl, and R' may be vinyl (CH₂=CH-) or ethyl. The eight commonest naturally occurring alkaloids compriso four structurally identical stereoisemeric pairs.

(1) Cinchenine and cinchonidine,

 $C_{10}H_{22}ON_2$ (R is H-; R' is $CH_2=CH-$).

(2) Hydrecinchonine and hydrecinchenidine, C₁₉H₂₄ON₂ (R is H-; R' is CH₃-CH₂-).

(3) Quinine and quinidine,

 $C_{20}H_{24}O_2N_2$ (R is CH_3O_- ; R' is $CH_2=CH_-$).

(4) Hydrequinine and hydrequinidine,

$$C_{20}H_{26}O_2N_2$$
 (R is CH_3O_- ; R' is CH_3-CH_2).

To this list may be added three other alkaleids belonging to the same chemical group. These have a phenolic character, and have not a methexyl but a hydroxyl group in position 6 of the quineline nucleus. Cupreine,

C₁₉H₂₂O₂N₂ (R = HO-; R' = CH₂=CH-), was discovered in euprea bark derived from Remijia prdunculata (Paul and Cownley, Pharm. J. 1881, 12, 497; 1884, 15, 221, 401; Hesse, Ber. 1882, 15, 854; Annalen, 1884, 225, 95), but is now unobtainable. It is the phenol corresponding to quinine, for it yields the latter on methylation (Grimaux and Arnaud, Compt. rend. 1891, 112, 774). The other two, hydrocupreino and hydrocupreidine,

C₁₉H₂₄O₂N₂ (R = HO-; R' = CH₃-CH₂-), are not knewn to occur naturally, but can be obtained by the demethylation with mineral acids of hydroquinine and hydroquinidino respectively, or, in the case of the fermer, by the catalytic hydrogenation of cupreine (Giemsa and Halberkann, Ber. 1918, 51 1325). The phenol corresponding to quinidine, which would systematically receive the name cupreidine, is unknown, although Ludwiczakówna, Suszko, and Zwierzchowski (Rec. trav. chim. 1933, 52, 817) claim to have isolated it from the products obtained by the treatment of quinidine with sulphuric acid; their contention, however, has been convincingly refuted (Henry and Solemon,

J.C.S. 1934, 1923; Thron and Dirscherl, Annalen | Thus Claus and Weller (Ber. 1881, 14, 1921)

1935, 515, 252). The above structural formula has not been Into above structural formula has not been automated to incommite and (quinonne-tindly elaborated for each allabid separately carboxyle acid) and not to its ethohorousic,
and follows mainly perhaps from work on Samap and vo Rosvall (Monatch. 1834, 15, 37)
conchounce The evidence for the skeletal prepared a series of yellow ethoholdes somene
structure of the other members of the groups with the ordinary white ethodoides by treating
proceeds in part from the stenity or simple not the free hases but the neutral hydrodides mutual relation of various degradation products; thus, for instance, cinchonine and cinehonidine give rise to identical products, however they are attacked [e q on oxidation there are formed cinchoninie acid (quinoline-4-carboxvise acid) and cincholosponic acid (3-carboxypipendine-4acetic acid)], the same applies, on the other hand, to quinine and quinidine, except that these give rise not to cinchoninic scid itself but to its 6 methoxy derivative, guminie acid Some evidence for the structure of certain of the alkaloids other than cinchonine arises also from the possibility of inter-ronversion among them; this, however, is very limited. Thus, ell the hydro aikaloids (R'=CH3-CH3-) are obtainable from the corresponding vinyl bases by catalytic hydrogenation Again, cinchonidine was obtained by Koenigs and Husmann (Ber. 1896, 29, 2185) from cinchonine by the action 1890, 29, 2489) from Charponne by the action provides party from the task that the season and of anyl elochole potassium phydroxide, and this can be demethylated (e.g. by he string with kind of interconversion was later extended by hydrochlorno acid), there are then formed Rabe and his pupil (Annelse, 1932, 492, 242) phenole haves and methyl chloride (see, 29, to the whole ceres Finally, in evidence of the llesse, Annalen, 1880, 205, 314). The other Rabe and his pupils (Annsien, 1902, 492, 242) to the whole series Finally, in evidence of the above list) of the cinchona alkaloids, may be mentioned the fact that each such pair gives rise to three transformation products, the identity of which is the same whichever member of the pair forms the source. These three products are, moreover, either actually isomeric with the parent alkaloids, or, at all events, have acid), quinine and quinidine furnish the corre-the full skeletal structure of the parent alkaloids aponding 6-methoxyquinoline derivatives. Thus preserved intact. They are (see below) the Skraup (Nonth, 1881, 5,837) found that quinne "icines" (toxines) of Pasteur and of ron Miller on oxidation with chromio acid gave riso to and Robdy, the ambydro-bases (ninchene, quinnes) of Comstock and Koenigs, and the ketones acid, and that this, on more vigorous oxidation (cinchoninone, quimnone) of Rabe. Thus, with the same resgent, was degraded to the emchonino and cinchonidine give use to one same pyridinetricarboxylic acid which could and the same cinchonicine (cinchotoxine), to be obtained in like manner from cinchonino one and the same cinchene, which only differs send, the exidation product of cinchenine from the parent bases by the elements of water ,

and to one and the same cinchoninone, differing from the parent bases by two hydrogen atoms. The evidence for the above structural formula

is in brief as follows.

Both nitrogen atoms in the molecule are tertiary and capable of salt formation. Thus not only are both neutral and seed salts known, such as B HCl, B H, SO, and B 2HCl, B H, SO, but also mixed acid salts of the type B, 2HCl H, SO, (Gramaux, Compt rend. 1892, 115, 608)

The alkaloids combine with one and with two molecules of an alkyl habde, forming monoand dislky halides, which have the characteristic properties of quaternary ammonium salts The nitrogen stom in the quinuclidine half of the molecule is more strongly hasic than that in the quinoline half and is the one involved in nentral salt formation ; it is at the quinucledine nitro- The constitution of the latter said being known gen atom also that the sikyl and halogen groups (see Skraup, Annalen, 1890, 201, 291), it followed are added in the ordinary monoalkylhahdes, that the methoxyl group in quininic acid was

showed that einchonding ethohromide was exidused to cinchoninic acid (quinoline-iwith ethyl iodide ; both forms furnished one and the same diethiodide when treated with a further molecule of ethyl iodide. Skrauplater (shid. p. 433) oxidised the yellow form of emchonine ethiodide and obtained the ethiodide of emchanine send, thus showing that in emchanine by driedide it is the quinuchdine basis centre which is satisfied. Similarly, Claus and Mallmann (Ber. 1881, 14, 76) found that the treatment of quinine methodide with ethyl sodide gave rise to a different product from that obtained by the action of methyl iodide on quanto ethiodide. The two "ethmethiodides" were isometic (cf. Claus, Annalen, 1892, 289, 232; and Konek, Chem. Zentr. 1936, u. 1332; 1937, 1, 1694 : 1938, 1, 751,

That one of the two oxygen atoms in quinne and quandine is a member of a methrxyl groop follows partly from the fact that these alkalands evidence for the presence of a met oxyl group, evidence, moreover, which acryes to indicate the position of this group in the molecule, arises from the fact that in those degradation experiments (such as oxidation) in which cinchonine and cinchonidine give rise to simple quinoline derivatives (such as cinchoninic

nucleus. He further showed (Monatsh. 1883, 4, 695) that quininic acid could be demethylated by means of hydrochlorie acid to the corresponding phenolic acid, to which he gave the name xanthoquininic acid, and that this, on decarboxylation, gave rise to the known 6-(or p-)bydroxyquinoline. There is, therefore, in quininie acid (as well as in quinine and quinidine) a methoxyl group in the 6-position of the quinoline nucleus, which is absent in the eases of einchoninic acid, cinchonine, and einehonidine.

The second oxygen atom in quinine and quinidine, or the sole oxygen atom in einchonine and cinchonidine, belongs to a bydroxyl group. This was first shown by Schützenberger (Compt. rend. 1858, 47, 233), who prepared the acetyl and benzoyl derivatives of einchonine and of quinine. Hesse (Annalen, 1880, 205, 314) has prepared the acetyl derivatives of all the four alkaloids. The demonstration, however, only acquires strict eogency when taken in conjunction with the later proof that these bases are ditertiary (see above). Independent of this proof, however, is the evidence adduced by Koenigs (Ber. 1880, 13, 285), who treated cinchonine hydrochloride with a mixture of phosphorus penta- and oxy- chloride and obtained a product, $C_{10}H_{21}NCI$, cinchonine chloride, in which the hydroxyl group had been replaced by chlorine. The naturo of the hydroxyl group was clucidated by Rabe and his collaborators (Ber. 1907, 40, 3655; 1908, 41, 62, 872; Annalen, 1909, 364, 330), who made the discovery that on very gentle oxidation with chromic acid einchonine lost two hydrogen atoms and was converted into the corresponding ketone, cinchoninone, being otherwise un-changed. Cinchoninone formed an oxime, and einchonine could be recovered again from its reduction products:

Ciuchoninone (R = H; R' = vinyl).

The cinchona alkaloids, therefore, contain a secondary alcoholic grouping, -CH(OH)-; its position in the molecule will be made clear in the sequel.

The existence of an unsaturated side-chain (i.e. of the vinyl group) in cinchonine, quinine, and their stereoisomerides is revealed by their behaviour with the halogens, the halogen hydracids, and with potassium permanganate. Laurent (Compt. rend. 1845, 20, 1587; Ann. Chim. 1848, (iii), 24, 302) first studied the action | of chlorine and bromine on cinchonine. Direct | addition of two atoms of the halogen occurs,

contained in the benzene and not in the pyridine | 17, 995; 1886, 19, 2853; 1892, 25, 1539) have also prepared these dihalogen addition compounds, as well as quinine dibromide, and made a detailed study of them. On treatment with alcoholic potash the dibromides lose two molecules of hydrogen bromide, dehydrocinchonine, C10H20ON2, and dehydroquinine, C20H22O2N2, being formed. These latter substances contain an acetylenic linkage in place of the double bond of the parent bases (R'= CH=C-) (Koenigs, Ber. 1895, 28, 1986). With the halogen hydracids, also, simple addition takes place, the side-chain R presumably becoming CH₃—CH(Hal)—. This cannot, however, as yet be considered as having been quite firmly established.

Zorn (J. pr. Chem. 1873, 8, 279), influenced by the contemporary researches of Matthiessen and Wright on narcotine and other opium alkaloids, was first led to study the action of strong hydrochloric acid on cinchonine and quinine (see also Hesse, Annalen, 1874, 174, 340; 1880, 205, 314). Skraup (Annalen, 1880, 201, 324) first treated cinchonine with hydrobromie acid, and Lippmann and Fleissner (Monatsh. 1891, 12, 327) heated quinine with hydriodic acid. In all these cases substances of the general typo HBHal,2HHal (where B stands for the original vinyl alkaloid) wero formed, except that the methoxylated alkaloids (e.g. quinine) were simultaneously demethylated. Coinstock and Koenigs (Ber. 1887, 20, 2510) showed that the reaction proceeded not only on heating but also at room temperature, and obtained in this way not only the ordinary ehloro. and bromohydrocinehonine but also eliloro. and bromoliydroquinine, which still contain the methoxyl group. A copious literature has accumulated dealing with this reaction,

The formation of halogen derivatives, whether by the addition of the free elements or by treatment with the hydracids, indicates only the presence in the molecule of an unsaturated. centre. The existence of a side-chain, and the fact that the double bond is contained therein, is shown by the behaviour of the vinyl-alkaloids with potassium permanganate. Whereas the hydro-alkaloids react but slowly or not at all (it is this property which led to their discovery and isolation), the unsaturated alkaloids instantly decolouriso a dilute, ice-cold, acid solution of this reagent. Kerner first studied the behaviour of quinine (Zeit. f. Chemic, 1869, 5, 593) and found that formic acid was produced together with a substance which he called quitenine, C₁₉H₂₂O₄N₂ (see also Skraup, Annalen, 1879, 199, 348). Similarly, Willin and Caventon (Annalen, 1870, Suppl. VII, 249), Hesse (Annalen, 1875, 176, 232), and Skraup (ibid. 1879, 197, 376) obtained cinehotenine, C₁₈H₂₀O₃N₂, from cinchonine, Skraup and Vortmann (ibid. p. 235) obtained einchotenidine from einchonidine, and Forst and Bochringer (Ber. 1882, 15, 1659) oxidised quinidino to quitenidine. These "tenines" result by the loss of one earbon atom as formic acid; they are, unlike the parent alkaloids, saturated substances which, for example, do not react cinchonine dichloride and dibromide being with hydriodic acid; they are tertiary bases formed. Comstock and Koenigs (Ber. 1881, and still contain a hydroxyl group; and finally

they are carboxylic acids which form salts and esters. These results were summarised by Skraup (Ber. 1895, 28, 12), who justifiably drew the conclusion that the unsaturated cinchons alkaloids contain a vinyl group, and that the "tenines" are the otherwise unchanged carboxylic acids (R' = -COOH).

Evidence to the same effect has been obtained by Seekles (Rec. trav. chim 1923, 42, 69); under the influence of ozona tha cinchona alkaloids form ozonides, which on hydrolysis give rise to aldehydes (R'=-CHO), correspondrise to indendues (N = CHO), corresponding to the 'tennes.' Seekles gave these the termination al, thus enchoninal, C₃, H₃,O₄N₂, and quinnal, C₃, H₃,O₅N₂. The position of the vinyl (or ethyl) group in the molecule will

appear in the subsequent discussion

The skeleton of the molecule of the emchons alkaloids naturally and historically falls into two portions or "halves", the quinoline portion or tha "first half," and the remainder of the molecule or the "second half," The molecule readily undergoes fission in a corre-sponding manner, there being formed on the one hand simple derivatives of quinoline or 6-methoxy quinoline, and, on the other hand, more complex derivatives of piperidine. The structure of the skeleton, and at the same time the position of the hydroxyl and vinyl (or ethyl) groups, was deduced from a study of such degradation products
There are, broadly, three ways in which such a

fission can be effected.

The first method, due mainly to Skranp, is by exidation with chromic acid, when the if first half " gives rise to the quinolinecarboxy he acids already mentioned, viz cinchonime and quininic acids. They are aparingly soluble readily seolable substances, and it is in their filtrates and mother bouors (which when simply evaporated afford intractable syrups) that the following substances derived from the "second half" are found meroquinens (from the ving I alkaloids), cincholoipon (from the hydroalkaloids), and the oxidation products of these, incholorponic and leiponic seids (Skraup, Monatsh 1853, 9, 783, 1896, 17, 365, Koenigs, Her. 1894, 27, 1501).

Meroquinene (R'1s CH2 - CH-) Cinchalofpon (R* is CH - CH -).

Cinchololponic acid.

Loiponic acid.

Exactly the same substances result from the bandation of the three sets of transformation products referred to above, ex. from the "toxines" (Skraup and Würstl, Monatsh. 1839, 10, 220), the anhydro bases (Comstock and Koengs, Ber 1884, 17, 1990), and from Rabe's Letones (Ber. 1907, 40, 3657), thus showmg the close relationship of these substances to the parent alkaloids. The same applies also to the saturated carboxylic acids, the "tenines" (formed by oxidation of the parent vinyl alkaloids with potassium permanganate), except that in this case neither meroquineno nor cincholoipon are formed, but only the quinoline-earboxylic acids, and cinclioloiponic and loiponic acids (Skraup, Monatsh 1889, 10,

The second method of fission was discovered by Koenigs. When the hydrochlorides of the cinchona alkaloids in chloroform suspension sre treated with phosphorus pentachlorido there are formed, as olready mentioned, substances such as emchonine chloride and quinino chloride, in which the original hydroxyl group is replaced by chlorine (Comstock and Koenigs, Ber. 1884, 17, 1984, 1885, 18, 1210. For the correspond-ing broundes, see C. F. Bochringer und Secher, C.m. b. II., Rabe, Cohaure, and Scheel, G.P. 992340 and 592341, 1834). These chlorides, on treatment with alcohole potash, lose a moleculo of hydrogen chloride with the production of a double bond. The "anhydro-basea" so formed are known as quinene, cinchene, hydroquinena (Giemsa and Halberkann, Ber. 1921,54,[B],1194) and hydrocanchene, each being the identical product from two stereosomeric parent alkaloids, thus both quining chlorids and quinidine chloride give rise to one and the same quinene.

Quinene (R is CH₂O--; R'is CH₂=CH--) Cinchene (R is H--: R'is CH. - CH-). Hydroquinene (Ris CH,O-; R'is CH,-CH,-) Hydrocinchene (R is H-; R' is CH,-CH,-)

When heated with hydrobromic acid, the anhydre-bases undergo an extraordinary reaction (Comstock and Koenigs, Ber. 1881, 14, 1854; 1885, 18, 1226, 2379; 1887, 20, 2674). The so-called apocinchene and apoquinene formed were found to be derivatives of phenylquinoline having the structure shown (see Koenigs, Ber. 1894, 27, 900; J. pr. Chem. 1900, 61, 1; Kenner and Statham, J.C.S. 1935, 299; and Kenner and Nandi, Ber. 1936, 69, 635). The vinyl group seems to be essential in the mechanism of this transformation, for, hydrocinchene does not react with hydrobromic acid (Koenigs, Ber. 1894, 27, 1504), and hydroquinene with hydrochlorie acid only undergoes demethylation (Giemsa and Halberkann, Ber. 1921, 54, 1195). This is suggestive in connection with the behaviour of the cinchona alkaloids with mineral acids (see below).

The reaction, howover, does not throw much light on the constitution of the alkaloids. Much more illuminating is the treatment of the anhydro-bases with phosphoric acid at 170° (Kocnigs, Bor. 1894, 27, 900), which constitutes the second method of fission of the cincliona alkaloids. Hydrolysis occurs, 2 molecules of water being taken up, and there are formed lepidine (4-methyl-quinoline) from cinchene, and its 6-methoxy-derivative from quinche. The former substance, derived from the "first half," is also formed from bydro-cinchene (Koenigs and Hoerlin, Ber. 1894, 27 2290). The "second half" gives rise to the same substances which are formed in Ekraup's chromic acid oxidation, viz. meroquinene from cinchene and quinene, and cincholoipon from hydrocinchene. The oxidation products of these, cincholoiponic and loiponic acid, are not formed in this reaction.

Before proceeding to the third method of fission, and to the remainder of the evidence for the structural formula set out above, it will be convenient first to deal with the constitution of the degradation products already mentioned, and to summarise the conclusions which can be drawn.

A full discussion of the nature of these substances, however, would go far beyond the

was early recognised, for Gerhardt (1843) bad obtained quinoline itself (then unknown) by the distillation of einebonine with potasb, and it was incidental to the investigation of these alkaloids that the constitution of this important heterocyclie base was studied. It was syntbesised almost simultaneously in 1879 by Koenigs, by Skraup, and by Baeyer.

In regard to the degradation products derived from the "second half," meroquinene, cincboloipon, eincholoiponic acid, and loiponic acid, it must suffice to append a bibliography of papers by Skraup and by Kocnigs, who, almost exclusively, studied and elucidated the constitution of these substances. [Skraup's papers are to he found in Monatsch., 1888, 9, 783; 1895, 16, 159; 1896, 17, 365; 1900, 21, 879; (with Piccoli), ibid. 1902, 23, 269. The relevant papers by Koenigs are in Ber., 1894, 27, 900, 1501; 1895, 28, 1896, 3150; 1897, 30, 1326; 1902, 35, 1349; 1904, 37, 3244; (with Bernhart), ibid. 1905, 38, 3049. For a very full account of the chemistry of meroquinene, see the experimental portion of Koenigs' "ehef d'œuvre" (Annalen, 1906, 347, 193); the theoretical portion of this paper (p. 143 et seq.) is entirely devoted to a full and valuable review of the then existing knowledge of the constitution of the cinchona alkaloids. For a synthesis of cincholoiponic acid, see Wohl and Losanitsch (Ber. 1907, 40, 4698; Wohl and Maag, ibid., 1909, 42, 627).]

The molecules of cinchoninic acid ("first

half") and of meroquinene ("second half") account between them for all the 19 carbon atoms of einchonine; further, the constitution of meroquinene fixes the vinyl group in a 3position to the piperidine nitrogen atom; of the three such possible positions, two are identical, structurally being only stereochemically distinct, while the third, made improbable by the manner of formation of meroquinene by Skraup's method of oxidation, is impossible in the light of its formation by the hydrolytic process of Koenigs. Finally, seeing that the nitrogen atom in the "second half" of the cinchonine molecule is tertiary, while that of meroquinene is secondary, and because no further carbon atoms (such as a N-methyl group) can be accommodated, it follows that either the acetic acid residue in the 4-position of meroquinene or the carbon atom of the earboxyl group of cinchoninic acid, must, in some way, be joined to the nitrogen atom in the einchonine molecule. Of the several possibilities which these considerations open for the structural formula of cinchonine, I and II (p. 142) were envisaged by von Miller and Rohde (Ber. 1895, 28, 1059); II which was originally preferred, was later rejected in favour of I by Koenigs (J. pr. Chem. 1900, [ii], 61, 11): be proposed the name "quinuclidine" for the parent structure, from which the "second half" of the following formula is derived hypothetically, and succeeded in preparing two stereoisomeric 3-ethylquinuclidines, one from meroquinene, and the other, by a different method, from eincholoircope of this article. It will suffice, in the case pon (Ber. 1904, 37, 3244). The formula of the quinoline derivatives (from the "first accepted at the present time, which is set forth half") to mention, as a matter of historical at the beginning of this section, was first coninterest, that the presence of a quinoline nucleus sidered by Rabe and Ritter (Annalen, 1906,

Cinchonine (von Miller and Rohde, 1895, I. Koenigs-

Cinchenine (von Miller and Rollde, 1895, 11)

350, 180); it was adopted by Koenigs, Bernhart, and Ibele (Ber 1907, 40, 648, 2873), hy Rohde and Antonaz (161d. 1907, 40, 2329), and finally aingled out by a process of elimination by Rabe, Kuliga, and Naumann (Annalen, 1909, 365, 353). And it is this work of Rabe's which must now bring us back to the consideration of the third method available for the degradation of the cinchona alkaloida.

obtained not only from quinine but also from quinidine. These substances, which are now. for reasons to be stated below, perhaps best described as the einchonatoxines or briefly as the torines, have the constitution shown above. The study of the toxines has played an essential part in the investigation of the parent alkaloids, but the manner of arriving at the constitution of the former, which can hardly be dealt with apart from a description of the researches which led to the formula of the latter Nevertheless, will not be indicated in detail

some salient points of interest may be mentioned.

I'wo other methods exist whereby the cinchons alkaloids can be converted into the isomeric toxines. One of these was unwittingly dis-covered by Claus and his pupils in 1878, and the other was developed by von Miller and Bohde in 1894 while engaged on a study of the Claus reaction The former chemist (Ber. 1878, 11, 1820) found that the quaternary addition compounds of the cinchena alkaloids with one molecule of an alkyl halide, when boiled with potassium hydroxide in aqueous solution, reverted to tertiary bases. He regarded these as ordinary homologues of the parent alkaloids, many substances of the general type " methyl cinchonino" were described in sovemi papers (see, for instance, Claus and Muller, Ber 13, 2200, Claus and Malmann, bid 1881, 14, 76, Claus, Annalen, 1804, 269, 232) von Miller and Rohde (Ber 1804, 27, 1187) found that thee "homologues" formed phenylhydrazones, and could not therefore have the parent atructure The parent alkaloids were not attacked by the reagent used (a solution of phenylhydrazine in accure acrd) unless heated with it at 100° for a long time. It was then found (ibid p 1279) that einchonine on heating with dilute acctic acid at 105° for 24 hours was transformed into a ketonic base, which reacted with phenyl-hydrazineatonce, and which, on methylation with methyl todide and sodium hydroxide, afforded a N-methyl derivative identical with Claus's "methyleinchonine" The Actenie base, on account of its alleged possonous properties and loss of antipyretic action, was given the name emchotoxine " (161d. 1895, 28, 1058) Its identity with Pasteur's cinchonicine was also auggested. Apart from their physiological properties, the toxines were found to be remarkable in several other ways As Pasteur had already noted, they had a greatly diminished optical rotatory power, and the toxines from stereo-isomeric pairs of parent alkaloids of opposite eign of rotation were identical; the alkaloids had become converted into ketones and secondary bases; finally, the rupture of the C-N linkage was remarkable in taking place more readily with weak organic acids (acetic, tartaric, phosphorie) than with strong mineral acids In 1833 Paster (Compt reed. 1853, 27, 110) auch as hydrochloric acid (labe and Mellian, discovered that the acid adiphates of the Ber 1910, 43, 3309; Rabe, 1504, 1912, 48, canchona alkalouds melt when heated with a [2927] Diddle (bld 1912, 48, 520, 2832) little water at 140° and give use to bomendes and adopted the same observation compared to the compared of the compared to the cause of the rection, and in the which he called einchonicine, and from quantum course of detailed physics chemical researches quincine. Moreover, einchondine gave rise with several pupile (J. Amer. Chem. Soc. 1913, to the same einchonicine which was also formed 35, 418; 11915, 37, 2003, 2003; 1916, from einchonnic. Similarly quincina was 23, 201; 1917, 39, 968) reached the ronions, nor by the anions of acids, but by undissociated molecules of acids, a sufficient concentration of which was only attainable in the case of weak organic acids. Rabe (Ber. 1907, 40, 3282; Annalen, 1909, 365, 366; Rabe, Schneider and Braasch, ibid. p. 377) pointed to the analogy between the toxinetransformation and the conversion of narcotine into nornarceine. Both cinchonino and nareotiae are 1:2-amino-alcohols, and Rabe adduced other illustrations in support of the general view that such substances are liable to undergo the intramolecular rearrangement which underlies the conversion of cinchonine into cinchotoxine, and it has recently been suggested (Konopnicki and Suszko, Bull. Acad. Polonaise, 1929, A. 340) that the conversion proceeds in two stages:

Substances of the intermediate enolic stage, in which the enolie group is fixed by intramolecular participation in an ether function, have been isolated, and it is proposed that the von Miller and Rohde toxine nomenelaturo be reserved for the final ketonic substances, while l'asteur's original ".ieine" nomenclaturo be adopted for the new enol-ethers. The toxine transformation can easily be brought into excellent accord with modern electronic theory (sec, for lustance, Hanhart and Ingold, J.C.S. 1927, 997).

The toxines react with hydroxylamine to form ketoxines and condense with one molecule of amyl nitrite in the presence of sodium ethoxide giving rise to isonitrosoketones of the type

When Koenigs proposed his formula for einchonine in 1900 lie stated also that einchotoxine must in conformity therewith have the structure indicated below.

Cinchotoxine (Koenigs, 1900).

Such a substance should react with two molecules of amyl nitrite to form a dissonitrosoketone of the type

clusion that it was catalysed neither by hydrogen | von Miller, Rohde, Brunner and Fussenegger (Ber. 1900, 33, 3214) failed to make einchotoxine react with more than one molecule of amyl nitrite. The final disproof of Koenigs' formula, and support for the formula proposed by Rabe, came on the one hand from Koenigs himself and on the other from Rabe. In each case oximino- or isonitrosotoxines were shown to undergo smooth rupture, into simple quinolino derivatives, from the "first half," and piperidine derivatives from the second, when subjected to the Beckmann rearrangement. Thus, Koenigs, Bernhart and Ibelo (Ber. 1907, 40, 648, 2873) treated the oxime of N-methyleinehotoxino with phosphorus pentachloride, and hydrolysed the product with aqueous-methyl alcoholic sodium hydroxide, or with diluto hydrochloric acid. There were formed 4-aminoquinoline, einchoninic acid, N-methylmero-quineno and (probably) β -4-(N-methyl-3-vinylpiperidyl)-ethylamine, products which were decisive for Rabe's formula as against that of Koenigs. Again, Rabe (with Ritter, Ber. 1905, 38, 2770; with Ackermann, ibid. 1907, 40, 2013) subjected isonitroso-N-methyleineliotoxino and isonitroso. N-methylhydroeinehotoxino to the Beekmann rearrangement, and obtained einehoninic acid and, from the first, tho readily isolable nitrile of N-methylmero-quinene, from the second, the corresponding eincholoipon derivative. At about this time, also, as has already been mentioned, Rabe succeeded in oxidising the secondary hydroxylic cinehona alkaloids to the corresponding ketones, in itself decisive evidence for his secondary hydroxylic formula as against Koenigs' tertiary one. As he pointed out, however, other formulæ, also containing a secondary hydroxyl group, were conceivable, and these were finally disposed of by studying the action of amyl nitrito on the ketones (Rabe, Ber. 1908, 41, 62; Rabe, Kuliga and Naumann, Annalen, 1909, 365, 353). The alternativo formulæ envisaged by Rabe contain a -CH2-COgroup, and demand the formation of isonitrosoketones. Instead, it was found that fission once again took place in accordance with the scheme

There were formed, that is to say, einchoninie acid and oximinovinylquinuclidine.

Oximinovinylquiauclidine.

Quininone, similarly, gave rise to this oximinosubstance and to quininic acid, and hydroeinchoninine furnished cinchoninie acid and oximinoethylquinnelidine. The oximino- sub-stances on hydrolysis, afforded hydroxylamine, with meroquinene from the one and cincholoipon from the other.

The constitution of the cinchens all aloids as well as of the cinchonatoxines had, in this way,

been finally established.

(b) Synthesis .- This constitution was venified in 1931 by the synthesis of hydrogumme and hydroquinidine (Rabe, Huntenhurg, Schultze and Velger, Ber. 1931, 64, 2487). The synthesis is in four phases:

Synthesis of quininic acid.

Synthesis of homocincheleipon (VI)-

Condensation of quininic acid with homo-eincholoipon to form hydroquinetoxine. Conversion of hydroquinetoxine to hydro-

quinine and hydroquinidine. Synthems of Quininic Acid. The first phase, which constitutes a modification of Knorr's p. 1343) quineline synthesis (Annalen, 1886, 236, 69), proceeds in aix stages ;

I. Acetoacetie ester is condensed with panisidine by heating the components tegether

2. The resulting p methoxyacetoacetanilide (I) is ring closed by means of 90% sulphune seid to 2-hydroxy-6 methoxy-4-methylquinoline (III).

3. This is converted to the corresponding chloromethoxymethylquinoline by treatment with a mixture of phosphorus oxy- and pentachloride.

4. The chlorine stom is then chminsted by reduction either catalytially in alkalino alcoholie solution, or by means of aluminium and acetic acid. There is then formed p methoxylepidine (6 methoxy-4-methylqumoline)

The direct oxidation of 6-methexylepidine does not proceed smoothly, and it was found better-

finally-

6. To exidise this styry lquinoline derivative with cold potassium permaneanate in acucous pyridine solution to 6-methexyquinoline-4carbovyle acid (quinme acid); cf. Kaufmann and Peyer, Ber. 1912, 45, 1805.

Synthesis of homoCincholospon -The second phase, the synthesis of homocincholoipon, proceeds in three stages. The method used had been worked out for a lower homologue by Rabe and Kindler (Ber. 1919, 52, 1846). starting material, B-collidine, or 4-methyl 3. ethylpyridine, which was discovered by Greville Williams (1855) and Orchener de Connek (Compt. rend. 1880, 91, 296; Ann. Chim. 1882, [1] 27, 469) in the products of distillation of emehonine with potash, had been synthesised by Ruzicka and Fermair (Helv. Chim. Acta, 1919, 2, 338; see also Rabe and Jantzen, Ber. 1921, 54, 925; E. Königs and Ottmann, shid.

1. B Collidane at 85° is condensed by means of

zine chloride with chloral.

2. The resulting compound (IV) is converted by means of alcoholic sodium ethoxide into the acrylic acid derivative V, which is then-

3 Catalytically hydrogenated in hydrochlone acid solution to give homocincholorpon (VI)

The last stage results in a mixture of storeeisomerides of which homocinchololpon is one; it can be isolated by fractional crystallisation of the d-tartrates of their ethyl esters. The desired stereoisomeride is obtained in prependerating proportion only by rigidly adhering to a set of narrowly prescribed conditions in 5. To condense it with benzaldebyde in the carrying out the hydrogenation. Among these presence of zine chloride to the corresponding may be mentioned the recommendation that C-benzylidene compound (III), and then the platinum catalyst be specially activated by the method of Vavon (Ann. Chim. 1914, [ix]. 1, 149) and that the reduction be carried out at a pressure of 3 ntmospheres and at 60°-70°.

Condensation of Quininic Acid with homo-Cincholoipon.—In the next phase of the synthesis quininie ester is condensed with homoeincholorpon ester-the latter in the form of its N-benzoyl derivative-by the Claison method. Rabe and Kindler (Ber. 1918, 51, 1360) had already, thirteen years earlier, succeeded in condensing einchoninie ester with non-synthetic homoeineholoipon to form hydroeinehotoxine, and had obtained hydroquinotoxine from quinime ester in like manner (ibid. 1919, 52, 1842). The latter condensation, under the agency of sodium ethoride and in the absence of a solvent, was new repeated with synthetic N-benzoylhomocincholoipon ethyl ester. The product (VII) was hydrolysed by means of hydrochloric neid, the carbethoxy and benzoyl groups being removed, and hydroquinotoxine formed.

Conversion of Hydroquinotoxine to Hydroquinine and Hydroquinidine.—The conversion of hydroquinotoxine to hydroquinine and hydroquindine in the last phase of the synthesis, which proceeds in three stages, is the earliest phase to have been elaborated, and dates back to 1908. In the first two stages hydroquinotoxine is converted to hydroquininone, and this is then reduced.

Two methods exist for the conversion of einchonatoxines to einchonaketones. One was first described by Rabe in the case of einehotoxino (Ber. 1911, 44, 2058). The other was devised by Kanfmann and Huber (Ber. 1913, 46, 2913) for hydroemchotoxino and bydroquinotoxine. According to Rabe's method the toxino is bromunated by means of hypobromous acid to a N-bromotoxine, which then, under the agency of sodium ethoxide in alcoholic solution, undergoes intramolecular alkylation: bydrogen brounde is climinated, the quinuclidine ring is closed, and a cinchonaketone formed:

Knufmann's method is similar. By brominating with a mixture of bromine and strong hydrobromic acid a C-broinotoxine is formed:

The second stage is identical with Rabe's, a emelionnketone being produced also in this

ease by an analogous mechanism.

Each method has certain advantages over the other. While Rabe's procedure is applicable to both saturated and misaturated toxines, Kaufmann's reaction is unsuitable for the latter. Rabe's method, moreover, can be effected in one operation by the use of sodium hypobromite. On the other hand, the Cbromotoxines lend themselves to manipulation more readily than do the N-bromotoxines; they separate in the course of the bromination as finely erystalline dihydrobromides. For the complete synthesis in 1931, Rabe and his collaborators actually used Kaufmann's method.

The last stage, the reduction of hydroquininone to hydroquinme and hydroquinidine (for a discussion of the formation of both alkaloids from the one ketone, see next section (c)), was the first to be achieved, for Rabe, already in 1908, lind obtained cinclioning by reducing emchaninone with sodium in alcoholic solution, and with iron filings in acetic acid (Ber. 1908, 41, 67). Kaufmann and Huber (lc.) reduced hydrocinehoninone with palladium black and hydrogen and were the first to obtain two stereoisomeric earbinols: hydrocinchonine and hydroeinchonidine. The other hydro-alkaloids were prepared from non-synthetic sources in a similar manner (Rabe, Z. angew. Chem. 1913, 26 543). The vmyl alkaloids proved more difficult to prepare from the ketones, because the vinyl group on catalytic hydrogenation was also reduced. A reducing agent suitable for this purpose, which would attack the ketone group but kave the vinyl group and the quinoline nucleus intact, was finally found in aluminim metal, used on an alcoholic solution of the ketone in the presence of sodium ethoxido (Rabe and Kindler, Ber. 1918, 51, 466; Vereinigte Chiminfabriken, Zimmer & Co., G.m.b.H., G.P. 330813, 1920).

In the synthesis described in 1931, with which we are here concerned, the synthetic hydrogummone was reduced by means of hydrogen and palladium black. A freshly prepared solution of the erystalline ketone, which contains a preponderant proportion of one of the two tautomerie forms [see section (c)], gave largely hydroquinidine which was isolated as the tartrate. A little of a new stereosomeride (m.p. 121°; [a]n 4 70 9°) was also obtained. Hydroquimine was obtained by the reduction of an only equilibrium

mixture of tautomeric forms.

Although a complete synthesis of hydrocmchonine and hydrocinchonidine has not been formally described, these substances virtually be considered to have been synthesised. They have been obtained from hydrocinchoninone, and this from hydrocinchotoxine (Kaufmann and Huber, Ic.). Hydrocinchotoxine has been

at that time only been obtained by the osidation rubshols (the suslegues of the circhons sika-of N-benzoylhy drecirchotoxine (Kaufmann and leida), and the rubyl chlorides were prepared by one as we have seen, been synthesised. Finally, These substances are of purely candense unchonnine acid was synthesised by Pfixmper (J. pr. Chem. 1902 (n); 60, 201 ; f. Thoelgans are reparted their stereochemical (J. pr. Chem. 1933, 71, 387). The case is different for the unsaturated alkaloids enchannes are of the unsaturated alkaloids enchannes and looks entain four desimilar, asymmetric extensions and the unsaturated alkaloids enchannes are considered to the considered to t quinme, the ayntheses of which present peculiar difficulties.

It is of interest to note here, that hy an analogous series of reactions substances have been prepared from which the emchona alkaloids may be considered to be derived. In 1922 Rabe (Ber. 1922, 55, [B], 522) proposed the names "rubane" and "rubatoxane" for the following structures .

the cinchonatoxines and cinchonaketones respec | marised in Table II.

prepared from einchoninic ester and N-benzoyl- tively, were described by Rabe, Kindler, and homoencholopon ester (Rabe and Kindler, Wagner (Ber, 1922, 55, [B], 502); rubane itself Ber, 1918, 51, 1360). The latter substance had [in optically active d and l forms), the four

atoms, marked (1) to (4) in the following formula, in which Q at nds for the quinolyl or 6-methoxyquinolyl residue, and R' for the viuyi or ethyl group. Sixteen stereoisomeric forms (8 pairs of mirror-image enantiomorphs) of any one alkaloid should therefore be capable of existence. Cinchonine and cinchonidine are two such forms, but they are not an enantiomorphous pair.

Although the nitrogen atom is also unsymmetrical, it does not affect the number of possible stereoisomerides, for, when two saymmetric atoms are, directly or indirectly, linked together three times over, they constitute but one centre of dissymmetry, as can be seen with the aid of a model. This accounts, for example, for there being only two forms of camphor (Skraup, Ber. 1902, 35, 3981; Rabe, tbid. 1922,

55, 527). The The two problems which challenge the chemist are (1) to determine the relative configurations or "partial rotations" of these four atoms in the various alkaloids, and 2) to prepare, Rubatoxane. in each structural case, the 14 other possible Rubatoxanene and rubanene, the analogues of stereoisomerides. The present position is sum-

TABLE II

Case	Represented by the following alkabilds		" Partial Rotation "of asymmetric C atoms.			
		1	2	3	4	
1 2	Cinchonine, quinidine, and their dihydro derivatives em Cinchonine, ep(quinidine, and their dihydro-deriva	+	-	+	+	
3	tires epiCinchonidine, epiquinine, and their dihydro-	+	-	+	-	
	derivatives	+	_	-	+	
5*	Cinchonidine, quinine, and their dihydro-derivatives d Hydroquinine (d-hydrocinehonidine and the corre-	+	-		_	
-	sponding Vinyl alkaloids are unknown)	-	+	+	+	
6†	I llydroquinidino (I hydrocinehonine and the corre- sponding vinyl alkaloids are unknown)	_	+	_	_	
71 81	Not represented by known alkaleids	-	+	+	-	
81	Not represented by known alkaloids	_	+	_	+	

ensatlomorph of case 4 † enantiomorph of case 1.

following manner:

When einchonine ehloride in dilute sulphnrie acid solution is reduced by means of iron filings, the halogen atom is replaced by hydrogen, and there is formed "desoxyeinehonine" (Koenigs, Ber. 1895, 28, 3143; 1896, 29, 372). The same applies to the other naturally occurring alkaloids, a distinct desoxy-base being formed in each ease. These substances are of central significance for the solution of the problem, inasmuch as in their molecules earbon atom 4, but not 3, has become symmetrical. For this reason. Rabe and his collaborators (Annalen, 1932, 492, 242), mistrusting the validity of stereocliemical arguments based on substances obtained by means of phosphorus pentachloride, prepared the desoxy-bases by a different route, riz. by heating the hydrazones of the einchonaketones (c.g. cinchoninonehydrazone) with powdered potash:

$$C \sim N - NH_2 \rightarrow CH_2 + N_2$$

a method due to Kishner (J. Russ. Phys. Chem. Soc. 1911, 43, 1398, and later papers). modification of the argument, however, was

necessary.

The einchona-toxines and -ketones furnish eases in which both earbon atoms 3 and 4 are rendered inactive together, while carbon atoms I and 2 remain unchanged. It is this circumstance which accounts for there being only one toxine and only one ketono for any one given pair of stereoisomerie parent alkaloids. The ketones are at first sight, like the desexybases, symmetrical about earbon atom 4 but not about earbon atom 3. Rabe showed, however, that while this may be the easo for the rolid substances, in solution enolisation occurred, and dynamic equilibrium was established between the keto- and the enol- forms: the einchonakctones are tantomeric (Rabe, Kuliga, Marsehall, Naumann and Russell, Annalen, 1910, 373, 85). Thus the ketones are slightly soluble in strong caustic alkali solution; they form O-benzoyl derivatives and they exhibit mutarotation. The asymmetry of centre 3 being destroyed in the enol- form, complete racemisation of this centre, in the keto- form also, must take place in solution.

That the configuration of centres I and 2 is the same for all the alkaloids was foreshadowed by Pasteur (Compt. rend. 1853, 37, 110), who observed that any one pair of stereoisomerie alkaloids was transformed to one and the same feebly dextrorotatory "-ieine" (toxine). Howard (J.C.S. 1873, 26, 1177) also noted that the optical rotatory powers of cinelionieine and of quinicine were, fairly accurately, the arithmetic means of those of cinchonine and cinchonidine, and of quinine and quinidine respectively. The extention of this argument from one stereoisomerie tion products of the einchonaketones. pair to another structurally distinct pair is made possible for all the vinyl alkaloids on the one hand, and for all the hydro-alkaloids on the other, by the fact that they all give rise to one and the same optically identical meroquinene or cincholoipon, as the case may be. These degradation products of the "second half"

This information has been obtained in the stereochemical gap between the saturated and unsaturated series was first closed by Koenigs (Ber. 1902, 35, 1350), who reduced meroquinene by means of zine dust and cold, fuming hydriodic acid. and obtained eincholoipon, optically identical with that gained from the bydroalkaloids by degradation. Since that time the transition from the vinyl to the hydro-alkaloids has been often effected by the method of catalytic hydrogenation.

Directly or indirectly, then, all the einehona alkaloids give rise to one and the same eineholoipon, and so have the same configuration about centres 1 and 2, the total contribution of which is weakly dextrorotatory, as shown by the optical rotatory power of the ketones and the toxines. The individual configuration at each of these centres has been worked out recently (Henry, Solomon, and Gibbs, J.C.S. 1937, 594; 1938, 6) by a study of the isomerides of the unsaturated alkaloids formed by the action of mineral acids (see section (d) below), and the results are included in Table II. For the present purpose it will suffice to regard carbon atoms I and 2 as constituting a single asymmetric centre, which is dextrorotatory in total effect in the naturally-occurring alkaloids. On this basis there will be 8 stereoisomerides in any one case of a single structural type. In the hydroquinine series, 6 of these 8 cases are represented. these, two, l-hydroquinine and d-hydroquinidino, occur naturally; their full mirror image enantiomorphs, d-hydroquinine and l-hydroquinidine have been synthesised by Rabe and Schultze (Ber. 1933, 66, [B], 120); the remaining two, epihydroquinine and epihydroquinidine, wero prepared from hydroquininone by reduction, as well as from hydroquinine or hydroquinidino by treatment with anyl alcoholic potash (Rabe and others, Annalen, 1932, 492, 242).

In spito of identity of configuration at centres and 2, the stereoisomerides einehonine and einehonidine-to take an instance-are, nevertheless, distinct, and must therefore havo opposite configurations either about carbon atom 3, or about carbon atom 4, or at both these centres. Now, the distinction between the two is preserved in the desoxy-bases; desoxyeinchonine is not the same as desoxycinchonidine. although they are identical at centres 1 and 2 and also at 4, being symmetrical at the latter point. The two desoxy-bases are therefore epimeric about carbon atom 3. The same applies to the parent alkaloids, except that these may or may not be epimeric also about carbon atom 4. Tho ideal way to settle this last point would be by an analogous procedure. Unfortunately, however, no derivative of the einchona alkaloids is known in which earbon atom 3 becomes symmetrical while carbon atom 4 remains unchanged.

The problem is solved by studying the reduc-

Carbon atoms 3 and 4 in the ketones are racemised by the dynamic equilibrium of tantomeric forms. On reduction back to the secondary alcoholic condition, however, they should resume a fixed dissymmetry. Now, the production of two new asymmetric earbon atoms in a molecule already possessed of an independent contain the original centres I and 2 intact. The centre of dissymmetry, and optically active,

should result in four non-enantiomorphous position as epicinchonidine (- at 3; + at 4), stereoisomendes; these need not necessarily be This relation is illustrated below. formed in equivalent proportions. The reduction of the cinchona-ketones should therefore be expected to give rise to four stereoisomene carbinols; the original naturally occurring pair, and two others. This was first found to be the case with hydroeinchoninone (Vereinigte Chininfabriken, Zimmer & Co., G m.h H., G.P. 330813, 1920), which, on reduction with aluminium or zinc in alcoholic solution and in the presence of sodium ethoxide, furnished the presence of somain emousing interactions strongly destrordstory hydrocanchonane ([a]_b + 190°), the strongly Invortatory hydrocanchonation ([a]_b - 95°), and two other alkaloids, one of [a]_b +88.5° and the other of [a]n +48°. It was then pointed out by King and Palmer (I C.S 1922, 121, 2577) that, without having to use the questionable procedure of applying in a strictly quantitative sense Hudson's rule of optical superposition, it was nevertheless possible to deduce, justifiably enough, that the strongly dextrorotatory hydrocinchonine had a dextro configuration at 3 and also at 4, that similarly, the strongly levo rotatory hydrocinchonidine had a lavo con figuration at each of those centres; while of the two new alkaloids, both of a more feeble rotatory power than either of the preceding, naturally occurring stereoisomerides, one had a dextro- at 3 and a leve configuration at 4, and the other, rice rerez, was dextro at 4 and læroat 3 The argument was generalised by analogy to all the emehona alkaloids, but it was pointed out that, on the evidence, it was impossible to establish the constitution of the new " hydrocinehonines "

The justification for that generalisation, and the identification of each of the new "epi-bases" was furnished by Rabe, Irschick, Suszka, Muller, Nielsen, Kolbe, von Riegen and Hoch-stätter (Annalen, 1932, 492, 242) All the "epibases" (see Table 11) were prepared, on the one hand, by reduction of the ketones, as had been done in the Zimmer patent with bydrocinchonin one, and on the other hand, by epimerisation of the parent alkaloids by means of amyl not only to emchanidine (Koenigs and Husmann, Ber 1896, 29, 2185), but to a mixture in dynamic equilibrium of all four stereosomerides in question-einchonidine, epicinehonidine, epieinchonine, and unchanged einchonine It was found, in experimental verification of King and l'almer's analogical generalisation, that the two most feebly rotating anbstances in any one group of four sterrorsomerides were the new eni bases

Finally, all the new alkaloids were identified to the other. In the case of cinchonine, cinexample, one of the latter gave rise to desort, or serversomerace or index, only a problematic enchounce; it was hence identified as epithological enchounce; at 3, - at 4); the other famished desox curchounder, and was assigned its hydroquindine and decox curchounder, and was assigned its hydroquindine respectively, except that they

Dextro-configuration at 3.

Levo-configuration at 3 en Cinchonidane Cinchonidine (+628°: (-111°: - at 4) Desoxy canchonidane (4=symm.)

The synthesis of the optical aptipodes of hydroquinine and of hydroquinidine (Cases 5 and 6 in Table II) was effected by Rabe and Schultze (i.c.) by precisely the same method whereby, as already explained their naturally occurring stereoisomerides had been synthesised two years earlier.

The \$\beta\$ and y-carbon atoms of homocincho-lospon (which had then been synthesised, and formed one of the last stages in the main synthesis) are unsymmetrical, and constitute carbon atoms 1 and 2 respectively in the suh

alcohole potash. For, in the latter process, it sequent einchona molecule. homoCincholoipen was found, for example, that einchonine gave rise can therefore exist in two recemic modifications, and both were in fact formed. Of the resolved, optically active components of one of these forms, one, obtained by means of d tartane acid, has proved to be identical with d homocinchologyon (i.e. with that obtained by the oxidation of benzoylhy drocinchotoxine). The other, its levo-enantiomorph, obtained aimilarly by means of I tartane acid, was esternied and in the form of its N benzov | derivative, was now condensed with quininic ester exactly as before. Carbon atoms 1 and 2 in the resulting hydroby conversion to the chlorides, and reduction of quinotonine therefore were levorolatery in these to the decory-bases in each case. In each crown combined effect. This new hydroquinnoce as revue for incomplete the converted to a new hydroquinnoce as previously known desoxy-base, and the other before, which, on reduction, gave rise to a mixture of carbinols rontaining presumably four chonidine, and their two new congeners, for stereoisomerides. Of these, only d hydroquinine

had opposite signs of rotation. the rotations of the new enantiomorphs were respectively equal to those of hydroquinine and hydroquinidine. On admixture, each pair of enantiomorphis formed a racemic, optically

inactive compound.

The other stereoisomerie homoeineholoipons, also present in Rabe's crude synthetic product, were not isolated. They would form tho starting material for a synthesis of einchona alkaloids in which carbon atoms I and 2 would have configurations differing, relatively to one another, from that present in the natural bases. These other homocineholoipons would provide a route to synthesis of some of the isomerides, which enabled Henry, Solomon and Gibbs (l.c.) to propose separato configurations for enrbon atoms 1 and 2, and a discussion of which follows.

(d) Behaviour with Mineral Acids.—A broad distinction between the hydro-alkaloids and those of the unsaturated series is that the former are substances which are relatively stable and indifferent to the action of strong mineral acids. These reagents serve, at most, to demethylate the methoxylated hydro-alkaloids, hydroquinino and hydroquinidine, to hydroeupreino and hydrocupreidino respectively.

The unsaturated alkaloids, on the contrary, when treated with mineral acids, sometimes even under relatively mild conditions, undergo a series of changes the nature of which has been and, to a smaller extent, still is n vexed problem in the history of these nikaloids. The difficulty is to some extent an experimental one. The reaction usually gives rise not to a single substance, but to a variety of products which are difficult to separato; many have a marked tendency to form with one another loose compounds which behave as single substances. Some, particularly the phenolic bases derived from quinino and quinidine, are difficult to crystalliso and to characterise owing to instability in the air, or to indefiniteness of meltingpoint, or owing to the fact that they undergo slow decomposition on drying for analytical

This problem is still receiving active attention in several laboratories, but enough is now known to warrant the following tentative account, even though it may require modification in the near future. For this reason also, and in view of the complexity of the subject, it will be more convenient not to encumber the discussion with references either to particular eases or to the experimental evidence. The bibliography of the literature on which this account is based will be

found at the end of this section.

Three distinct types of substance may be produced when the unsaturated einchons alkaloids are treated with mineral acids. The first type (A) arises from the addition of the elements of the reagent to the double bond. The other two types result from the isomerisation of the parent alkaloids by (B) wandering of the double bond, and (C) loss of the double bond

Numerically sufficiently drastic; the phenolic bases apomorphs were quinine, apoquinidine, and their congeners are then formed. The three types have had assigned to them the following structural formulæ respectively, which are probably substantially correct:

$$\begin{array}{c} Q \\ \downarrow \\ CH(OH) & CH \\ \downarrow \\ H_2C & CH_2 \\ CH_2 & C \\ CH_2 & C \\ CH \\ CH \\ CD & H \end{array}$$

Type B.

Q stands for the quinolyl-, 6-hydroxy- or methoxyquinolyl residue.

A fourth type of substance is also formed. This ennuet, however, be considered here, as no general agreement has as yet been reached concerning its structure. This fourth type is represented by " δ -cinchonine" (" α - and β -cinchonlydrine") from cinchonine, "niquine" from quinine, and "niquidine" from quinidine.

Substances of typo A (X=halogen) are formed to the exclusion of the other types when any vinyl alkaloid is treated with very concentrated hot or cold halogen hydracid (e.g. hydrochloric acid, saturated at -17°). The same substances nre also produced when any of the transformation products, A (X = -OH), B, or C, are treated in similar fashion, except that a higher temperature is then requisite.

When the acids are dilute and hot (e.g. hydrochloric acid of sp.gr. I-125 at 140°), or when sulphuric acid (50% and upwards) is used instead of the halogen neids, all three types are formed side by side, except that in the ense by intramolecular ether formation involving the central hydroxyl group. In addition, the methoxylated alkaloids, quinine and quinidine, type C is absent. This statement nlso applies either to the parent alkaloids or undergo demethylation when the action is

No two parent alkaloids are known to give rise to any one and the same transformation product. It is assumed, therefore, that the configurations of the four asymmetric carbon atoms originally present are not affected by treatment with acids, and that the transformation products derived from other possible, but still unknown, parent types would similarly occur in distinct sets. The existence of three isoquinidines of one type renders this assumption untenable, at all events in the ease of type C. The third isoquinidine is not derived from quinidine, but probably from a stereoisomerido.

The following bibliography, on which the foregoing account of the behaviour of tho einehona alkaloids towards mineral acids is based, is arranged on a chronological basis of authors. The publications of any one author and his collaborators are not interspersed with those of other authors, so that the order of the entiro list is not chronological. Tho authors are listed in order of date of the earliest publication:

Zorn (J. pr. Chem. 1873, [ii], 8, 279); Hesse Zorn (J. pr. Chem. 1873, [1i], 8, 279); Hesse (Annalen, 1880, 205, 314; 1888, 243, 131; 1890, 260, 213; 1892, 267, 138; 1893, 276, 88, 125); Julius (Monatsh. 1884, 6, 750); Comstock and Koenigs (Ber. 1887, 20, 2510); Koenigs (Annalen, 1906, 347, 184); Jungfleisch and Léger (Compt. rend. 1887, 105, 1255; 1888, 106, 68, 357, 657, 1410; 1889, 108, 952; 1891, 113, 651; 1892, 114, 1192; 1893, 147, 42; 113, 651; 1692, 114, 1192; 1893, 117, 42; 1891, 118, 29, 536; 1894, 119, 1268; 1895, 120, 325; 1901, 132, 410, 828; Ann. Chim. 1920, 14, 59); Léger (Compt. rend. 1918, 166, 76, 469, 903; 1919, 168, 404; 1919, 169, 67, 797; J. Pharm. Chim. 1936, 23, 558); Lippmann and Fleissner (Ber. 1891, 24, 2827; 1893, 26, 2005; Monatsh. 1891, 12, 327; 1893, 14, 371; 1895, 48, 341; Sirroup, (Monatsh. 1891, 42, 431; Monatsh. 1891, 12, 327; 1893, 14, 371; 1893, 16, 34); Skraup (Monatsh. 1891, 12, 431; with Schubert, Monatsh. 1891, 12, 669; Ber. 1892, 25, 2909; Monatsh. 1893, 14, 428; 1897, 18, 411; 1899, 20, 571, 585; 1900, 21, 512; with Zwerger, Monatsh. 1900, 21, 535; 1902, 23, 455; 1904, 25, 894; Monatsh. 1901, 22, 171, 253, 1083, 1097; 1903, 24, 291, 311; with Eggrer Monatsh. 1903, 24, 660; Pure with Egerer, Monatsh. 1903, 24, 669); Pum (Monatsh. 1891, 12, 582; 1892, 13, 676; 1895, 16, 68); Neumann (Monatsh. 1892, 13, 651); Cordier von Löwenhaupt (Monatsh. 1898, 19, 461); von Arlt (Monatsh. 1899, 20, 425); Langer (Monatsh. 1901, 22, 151, 157); Havnička (Monatsh. 1901, 22, 191); Widmar (Monatsh. 1901, 22, 976); Kans (Monatsh. 1904, 25, 1145; 1905, 26, 119); Pfannl (Monatsh. 1911, 32, 241); Paneth (Monatsh. 1911, 32, 257); Bötteher and Horowitz (Monatsh. 1911, 32, 793; 1912, 33, 567); Rabe and Bötteher (Ber. 1917, 50, 127); Rosenmund and Kittler (Arch. Pharm. 1924, 262, 18); Giemsa and Bonath (Ber. 1925, 58, [B], 87); Fränkel and Buhlea (Ber. 1925, 58, [B], 87); Fränkel and Buhlea (Ber. 1925, 58, Frankel and Buhlea (Ber. 1925, 58, Frankel and Buhlea (Ber. 1925, 197 1925, 58, [B[, 559); Suszko et al. (Bull. Acad. Polonaise, A. 1925, 129; 1929, 340; 1933, 119; 1935, 65, 360, 415, 457, 465; Rocz. Chem. 1933, 13, 360, 464; Rec. trav. chim. 1933, 52, 839, 817; 1935, 54, 481; 1936, 55, 392); Ilenry (with Solomon, J.C.S. 1934, 1923; with Solomon and Gibbs, ibid. 1935, 966; 1937, 502); Carlon J.C.S. 1937, 1937, 1937, 1938, 19 592); Goodson (J.C.S. 1935, 1094); Solomon (ibid. 1938, 6).

6. Properties of Individual Substances. The following list, designed to give more detailed information concerning the discovery, occurrence, and properties of the individual alkaloids and their derivatives, is arranged in alphabetical order of the naturally occurring alkaloids (p. 128). After each description of an alkaloidal base there follow descriptions of its principal salts and derivatives, and tho more important of its transformation and degradation products.

Aricine (quinovatine), $C_{23}H_{26}O_4N_2$, named after the port of Arica in Peru, was discovered in Cusco bark (C. Pelletierana) by Pelletier and Corriol (J. Pharm. Chim. 1829, 15, 575). It forms large, colourless prisms from alcohol, m.p. 188° , $[a]_{D}^{15}$ -58.2° (c=2 in 97% alcohol), but +14.5° in acid solution. It is soluble at 15° in 100 parts of 90% alcohol, or 33 parts of ether, very soluble in coloroform and insolublo in water. With concentrated sulphurie acid it gives a greenish-yellow colour, changing to deep blue on addition of ammonium molybdate, and this becomes olive-green on warming. This colour reaction is also characteristic of cusconine. Nitric acid produces a deep green coloration, and affords a greenish-yellow solution. is astringent but not bitter in taste. Arieino It is a weak base, and its salts dissociate in water.

Thohydrochloride, B.HCl,2H2O. forms eolourless prisms, slightly soluble in cold water. The sulphate, B₂·H₂SO₄, consists of fine needles, fairly soluble in cold water. The acid sulphate, B·H₂SO₄, is less soluble. Aricino also forms an acetate, a citrate, a hydriodide, and two oxalates, the "binoxalate," B·C₂H₂O₄, 2H₂O, heing your springly soluble in cold water. being very sparingly soluble in cold water. Tho platinichloride, salicylate, and thiocyanato are also described (Hesse, Annalen, 1878, 185, 296; Moissan and Landrin, J. Pharm. Chim.

1890, 21, 337).

Chairamidine, C₂₂H₂₆O₄N₂, was isolated from the bark of *Remijia Purdicana* by Hesso (Annalen, 1884, 225, 253). It is accompanied in the bark by a series of other alkaloids, and is isolated by first removing these as sulphates, hydrochlorides or thioeyanates. For details of the method Hesse's paper should be consulted. Chairamidine is a white, amorphous powder containing 1H₂O. The anhydrous substance has m.p. 126°-128° (decomp.), and $[a]_D^{15} + 7.3^{\circ}$ (c=3 in 97% alcohol). It is insoluble in water, and readily soluble in alcohol, ether, ehloroform or benzene. It dissolves in concentrated sulphuric acid to a yellow solution, which slowly turns green. Animal charcoal removes it completely from a solution in acetic acid. Chairamidino is a weak base, and its solutions in alcohol are neutral to litmus. Its salts are amorphous.

Chairamlne, C₂₂H₂₆O₄N₂, also occurs in Remijia Purdicana (Hesse, Annalen, 1884, 225, 213), and is obtained as the hydrochloride from the mother-liquors remaining after removing tho concusconine as sulphate. The base crystallises with 1H2O from dilute alcohol (slender white needles) or from strong alcohol (thick prisms). The hydrate melts at 140°, and the anhydrous Hut in the case of type B, see footnote on page 150. base at 233°; [a]p about +100° in 97% alcohol.

dissolves to a colourless solution in concentrated sulphuric acid, but this slowly turns dark green. This colour is also produced when concentrated natric acad as added to a solution of chairsmine in acctic acid. In addition to the crystalline hydrochloride used for its separation, a sulphate, B, H, SO, 8H, O, is also known.

Cinchamidine. - See Hydrocinchonidine. Cinchona Febrifuge .- See Quinetum,

Cinchonamine, C1. H2. ON, was soluted by Arnaud (Compt. rend. 1881, 93, 593; 1883, 97, 174). It occurs in the bark of Remigia Purdicana together with cinchonine, concusconine, chair amine, chairamidine, conchairamine, and conchairamidine Hesse (Annales, 1884, 225, 211, 218) gives a scheme of separation. The emchon mine is purified by taking advantage of the involubility of its nitrate. The base crystallises from alcohol in needles or ortherhombie, tribeluminescent prisms, mp 181°-183° (Hesse), 193° (Armad), [a]¹³ +1211° (c=2 in 97%) alrohol) It is almost involuble in cold water, but slightly soluble in hot water, from which it one signify solution in the water, from which is crystallises in abort prisms Cinchonsmine dissolves in 31 6 parts of 90% alcohol at 17°, and in 100 parts of ether. It is also soluble in chloroform, benzene, light petroleum or carbon disulphide. Cinchonamine is said to be toxic, but more powerfully antipyretic than quinine Its solutions do not fluoresce, neither do they give the thallesoquin reaction. It differs from its isomerides. hadrocinehonine and bydrocin chonidine, in being readily attacked by potassium permanganate Canchonamine does not contain a methoxyl group, but forms a methodide and an aretyl derivative. With nitric acid it forms a yellow, amorphous, explosive dinstro denvative.

Cinchonamine is a strong discidic base which forms well-defined salts Of greatest interest is the nitrate, BHNO2 (minute prisms, mp 106°), the insolubility of which is found useful. as already stated, in purifying this alkaloid Arnaud (Ann. Clum. 1890, 19, 93) proposed the use of cinchonamine for the estimation of nitrates, but the method is limited in sta applica-The hydrochloride, BHCi, soft lamme, or B₂·H₂SO₄ (prisms), in contradistinction to a number of cinchons alkaleids, is very soluble in numeer of cincuous anagonus, is very sounds in water, as is also the acid aniphate, B H₂SO₄ (octahedra or prisms). The citrate, hydroboromide, hydrodide, melate, platmichloride, picrate, tartrate, and thiocyanate are also picrate, tartrate, and thiocyanate are also known. Boutroux and Genvresse (Compt rend. copper.

Cinchonicine .- See Quehotoxine under Cinchonine.

Cinchonidine, C1, H21ON2, occurs naturally in most cinclions barks, and particularly in C. succirulea It can also be obtained artificially by treating cinchonine with amyl alcoholic

It is soluble at 11° in 540 parts of 97% alcohol, 2185). It was first isolated from C. tucujensis but is readily soluble in other or chloroform. It by Winckler (Jahresbericht, 1847/1848, 620) by Winckler (Jahrebericht, 1847/1845, 620), who called it "quinidme." It was renamed "cinchondine" by Pasteur (J. Pharm. Chim. 1853, 23, 123), who suspected that the new alkaloid was isomeric with cinchonine. The confusion which resulted in the literature may be regarded as typical of the uncertainty, which existed in the middle of last century, concerning the adentity of several cinchons alkaloids (see e g. Hesse, Annalen, 1865, 135, 334: 1873. 168. 243; 1875, 176, 322; Ber. 1877, 19, 2149, 2152). The identity of cinchonidine was finally placed on a satisfactory basis by Skraup and Vertmann (Annalco, 1879, 197, 226), who also gave it the correct empirical formula. It is isolated (from the mother-liquors from quinine sulphate) as tartrate, and purified either as neutral sulphate (Hease, Annalen, 1880, 205, 196), or as acid sulphate; for the preparation of the latter, sulphate; for the preparation of the case is each gram of crude, dry enchondine base is dissolved in 21 cc. of 50% sulphure acid and 11 cc. of alcohol. The base is finally recovered and recrystallised from alcohol The product so obtained atill contains some quieine and hydrocinchonidine, which can be removed by reerystallisation from benzene, and fractional crystallisation from alrohol respectively (Buttle, Henry, and Trevan, Brochem. J. 1934, 28, 436).

Conchonidine crystallises in anhydrous, large trimetric prisms, or thin plates. Its meltingpoint has been given variously, ranging from 202° to 219 5°, the melting point given by Buttle, Henry, and Trevan (ibid, p. 438) for their pure circhonidino is 204 5°. It is lavorotatory · [o]₁^{17 a} ~107 9° (c=1 in 1 vol. alcohol+2 vols chloroform) (Lenz, Z. anal. Chem 1888, 27, 563); -86 2° (c=1-0955 ln there is 25, 27, 203); -50 2" (c=14935 in chloroform) (Rabe et al. Annalen, 1910, 373, 100), -111 0" (c=0 878 in 99% alrohol) (Rabe, 1c); -175" (c=0 735, ve. 3//40 solution in 91% H₅CO₂) (Buttle et al., te.). Cinchodidine is almost insoluble in water (1 in over 5,000); it dissolves in 300 parts of cold alrohol (sp.g., 0.935) or in 16 parts of 97% alrohol; in over 1,000 parts of dry ether, but in 188 parts of ether ap gr. 0-72. It is readily tion (Howard and Chick, J.S.C.I. 1909, 28, 53). soluble in chloroform or amyl alrohol. Cinchondine solutions neither fluoresce nor give the B HCi, H.O. transparent cubical erystals, is thallesoquin reaction (distinction from quining actuble in 200 parts of water at 27 (lloward and quantine). The alkaloid is distinguished and Perry, ibid 1905, 24, 128t) The sulphate, from its atcreoisomeride, einchonine, in being lavorotatory, in its greater solubility in ether, and by giving an insoluble tartrate. On oxidation with potassium permanganate it gives rise to cinchotenidine (see p. 154b), and with chromic acid it is degraded to the same aubstances that are formed from emchanine in like manner. On reduction with hydrogen in the presence of 1897, 125, 467) have described a scries of crystal- platinum or palladium catalysts, hydrocin-line double chlorides with cadmium, zinc and chonidine (q v) is formed. Cinchonidine gives hydrocmrise to the same cinchetexine (cinchenicine) that is formed from einchonine.

Canchonidine Sulphate, B. H.SO, crystallines from cold water in necelles containing BH,O, from het water in needles containing 3H,O, and from alcohol with 2H,O. The salt beromes anhydrous at 100° (m p. 205°, decomp), and repotash (Koenigs and Husmann, Ber. 1896, 29, absorbs 2H,O on exposure. At 15° the asit is soluble in 100 parts of water, 60 of alcohol, or has m.p. 166° (Neumann, Monatsh. 1892, 13, 1,000 of chloroform; at 25° it dissolves in 63 of 651).

parts of water or 72 of alcohol. The solubility in apoCinchonidine (Hesse, Léger), β-einchonidine water at 80° is threefold that at 25°.

Cinchonidine Acid Sulphate, B.H2SO4,5H2O, is readily soluble in water; $[a]_D^{15} -133.6^{\circ}$ (e= 0.93 of anhydrous salt, i.e. an M/40 solution, in water). The same rotation is observed in M/10solution, and in 0·1N-H₂SO₄. The tetra-sulphate, B·2H₂SO₄, H₂O, is slowly soluble in water.

Cinchonidine Hydrochloride, B.HCl, H2O, forms monoclinic prisms from water. It crystallises from alcohol in fine needles containing 2H2O. The anhydrous salt has m.p. 242°; $[a]_{0}^{20}$ -117.6° (c=1.214 in water). The salt is soluble at 12° in 30 parts of water, and in 300 parts of other; it dissolves readily in alcohol or in chloroform.

Tartrale, $B_2 \cdot C_4 H_6 O_6, 2H_2 O_6$ Cinchonidinc forms fine needles soluble in 1,265 parts of water at 10°, and almost insoluble in sodium potassium tartrato solution.

Other known salts of einchonidine include the dillydrochloride, hydrobromide, dillydrobromide, hydriodide, dihydriodide, nitrate, oxalate, acid tartrate, benzoate, salicylato and succinate. number of double compounds with metallic salts have been described.

Cinchonidine Methiodide, C19H22ON2 CH31, is formed from the components in alcoholic solution at room temperature (24 hours). Fine colourless needles, m.p. 248° (Claus and Bock, Ber. 1880, 13, 2192).

Acceptainchonidine, $C_{10}H_{21}(CO\cdot CH_3)ON_2$, results from the action of acetic anhydride at

60°-80°. Amorphous powder (Hesse, Annalen, 1880, 205, 319; Hilditch, J.C.S. 1911, 99, 238). a-Chlorohydrocinehonidine, better a-chlorodihydrocinehonidine (component of "hydrochlerocinchonidine "),

is obtained by the action of hydrochloric acid, either at room temperature with acid saturated at -17°, or at 85° with the ordinary concentrated acid. It is isolated from the products of the reaction as acid sulphato or tartrate, and then fractionated as dihydrobromide when it forms the more soluble component. The base (from alcohol) has m.p. 231° (decomp.), $[\alpha]_D^{23}$ -135.6° (c=0.5 in N-HCl) (Goodson, J.C.S. 1935, 1096).

The more sparingly soluble component in the dihydrobromide fractionation is a'-chlorohydrocinchonidine, which is epimeric with the aderivative at the carbon atom carrying the chlorine atom. The a'-derivative (base) crystallises from alcohol, has m.p. 246° and [a]_D²³ -62.5° (c=0.5 in N-HCl). Its acid sulphate, tartrate and dihydrobromide are also described (Goodson, ibid. p. 1097).

Iodohydrocinehonidine, better iododihydrocinchonidine (hydroiodocinchonidine),

$$C_{3}H_{6}N-CH(OH)-C_{7}H_{11}N-CH1-CH_{3}$$

apoCinchonidine (Hesse, Léger), β-einchonidine (Hesse, Neumann, Léger), y-zinchonidine, isocinchonidine (Hesse, Paneth),

$$C_9H_6N-CH(OH)-C_6H_{10}N>C=CH-CH_3$$

These four names have been used by the authors, whose names are bracketed after them, to designate a series of eight substances, the modes of formation and characters of which are assembled in the following table. Most probably only two pure substances, which are geometrical isomerides with the constitution indicated, are involved; some, if not all, of the eight varieties described (Table III, p. 154), may be mixtures in differing proportions.

The cinchonidine isomerides form monoacctyl derivatives and react with concentrated halogen hydracids to give the same halogenohydrociachonidines as are obtained from the

parent alkaloid.

Hydroxyhydrocinchonidine, better, hydroxydihydrocinchonidine,

$$C_9H_6N-CH(OH)-C_7H_{11}N-CH(OH)-CH_3$$

is formed together with β - and ap ocinchonidine in the sulphuric acid method of Léger (Bull. Soc. chim. 1919, [iv], 25, 575). Like these, it crystallises from dilute alcohol, but it forms a diacetylderivative. Hydroxyhydrocinchonidine m.p. $242^{\circ}-243^{\circ}$, and $[a]_{D}^{22}-101\cdot7^{\circ}$ in alcohol.

epiCinchonidine, $C_{19}H_{22}ON_2$, is epimeric with cinchonidine about the carbinol (CH·OH) group. It is obtained either by heating cinchonine with amyl alcoholic potash (300 g. alkaloid, 150 g. KOH, 5.5 litres amyl alcohol), or by reducing cinchoninone (100 g.) dissolved in a solution of sodium (84 g.) in alcohol (2,280 c.c.), with aluminium powder (84 g.). The epicinchonidine is isolated from the ether-soluble fraction of the sterco-isomerides formed, with the aid of tartarie, dihenzoyl- and dianisoyltartarie acids. It forms small, colourless plates from ether, m.p. $103^{\circ}-104^{\circ}$, $[a]_{D}^{20}$ (c=0.8044 in alcohol), is readily soluble in the usual organic selvents, and is non-fluorescent (Rabe and others, Annalca, 1932, 492, 253.)

Cinchoniding Chloride,

$$C_9H_6N$$
— $CHCI$ — $C_7H_{11}N$ — CH = CH_2 ,

is formed when anhydrous einchonidine hydrochloride (33 g.) dissolved in chloroform (300 c.c.) is slowly treated with phosphorus pentachloride 50 g.), care being taken to avoid any rise in temperature by cooling with water. The product crystallises from a mixture of ether and ligroin (b.p. 40°-50°) in white crystals, m.p. 108°, $[a]_{\rm D}^{13}$ +78.2° (c=2.020 in 99% alcohol); +90.9° (c=2.009 in ehloroform). The hydrochloride, C₁, H₂₁N₂Cl·HCl, has [a]_p²⁴ +24·16° (c=1.573 in water) (Comstock and Kernigs, Ber. 1884, 17, 1987; Pales Kuller Words II. 17, 1987; Rabe, Kuliga, Marschall, Naumann and Russell, Annalen, 1910, 373, 103). Desoxycinchonidine,

 $C_9H_6N-CH_2-C_7H_{11}N-CH=CH_2$

is obtained by heating einchonidine with hydrilis prepared from einchonidine chloride. A 5% odic acid (sp.gr. 1.7) on the water-bath. The base solution of the chloride (12 g.) in 10% sulphuric

TARLE III.

Name.	Manner of Formation from Cinchonidine.	Мър	[a]».	Author and Reference,
apoCinchonidino	Heating with 3 parts HCI (sp gr. 1-125) at 140°-150° for 6-10 hours.	225"	-129 2° in alcohol -160-4° in dil. HCl.	Hesse, Annalco, 1880, 205, 323
apoCinchonidine	Boiling with 4 parts of 50% H ₁ SO ₄ for 48 hours.	254 7 to 256 7°	-134 4° in alcohol166 6° in dil. HCl139 3° in slco-	Léger, Bull. Soc chim. 1919 [17], 25, 574.
β Cinchonidino	Heating with 3 parts HCI (sp gr. 1-125) at 140°-150° for 6-10 hours.	206° to 207°	-1814° in dil. HCl.	Hesse (l.c.).
β Ciuchonidine	From iodohydrocinchoni- dine and alcoholic KOH.	214	-171 5° in alcohol	Neumann, Mon atsh. 1892, 13 655.
β Cînchonslino	Boiling with 4 parts of 50% HaSO4 for 48 hours.	240° to 241°	-126 6° in alcohol, -181 5° in dil, HCL	Léger, Bull. Soc chim. 1919 [1v], 25, 573,
y-Cinchonidine	From iodohydrocanchoni- dane and alcoholic AgNO ₂ .	238°	-164 6° in alcohol.	Neumann, Blon- atsh, 1892, 13, 659.
140Cinchonidine	Solution in cold conc.	235°		Hesse, Annalen, 1888, 243, 149
isoCinchonidino	Heating with 6 vols. of 25 N. H. SO4 at 100° for 2 hours.	252°	-128° in alcohol- chiocoform.	Paneth, Mon- atsh 1911, 32, 259, 269.

has mp 60°-62°, and [o]_D¹³ -29 9° (c=2 006 in 99% alrohol), or -19 7° in chlocoform. On treatment with alcoholic potash it gives rise to cinchene (see under einehonme) (Koenigs, Ber. 1896, 29, 373; Rabe et al., Annalen, 1910, 873, 106). Cincholenidine.

C.H.N-CH(OH)-C,H,1N-CO,H, and formic acid are formed when cinchonidine is oxidised with potassium permanganate (Skraup and Vortmann, Annalen, 1879, 197, 235, Hesse, Ber. 1881, 14, 1892), It crystallises from alcohol in frathery crystals, or from water in monoclaric prisms containing 3H,O It has m p. 256° (decomp.), [a] -201 4° (e=5 m water containing 3 equivalents of HCI), is aparingly soluble in cold water, readily so in hot, and dissolves in 600 parts of boiling alcohol. It is soluble in acids as well as in caustic alkalis and is precipitated from the latter by earbon Cinchotenidine is non fluorescent.

treated repeatedly with alcohol or other, and the of alcohol at 17°, but in 28 parts of the boding insoluble einchonine either recrystallised from solvent; in 370 parts of cold ether (sp gr. 0-73).

acid is shaken with icon filings (17 g) for 72 boiling alcohol or converted into sulphate and hours. The product crystallises from legron, recrystallised from water. The enchemnes has m p 60°-62°, and [61] "-298" (=206) in filinally recrystallised from alrohol several time. to remove traces of quinidine. The alkaloid se prepared (adaptation of a method given by Hease, Annalen, 1862, 122, 227) contains about 15% of hydrocinchenine which can be removed hy fractional crystallisation of the dihydrobecomides, einehonine duby drohromide being the more soluble component (Buttle, Henry and Trevan, Biochem. J. 1934, 28, 437) or hy Thron and Duscherl's method (Annalen, 1935, 515, 252). Although the correct empirical formula was given by Laurent in 1848 (Ann. Chim. 1848, 19, 365), the formula, C₁₀H₁₄ON, proposed by Regnault in 1838 (Annalen, 1838, 28, 15) was current until 1879 when Skraup (Annalen, 1879, 197, 353) finally showed that Laurent's formula must be adopted. Cinchonine crystallises from alcohol or from ether in prisms, the recorded melting point is very vanable, and depends on the rate of heating (Hesse, Annalen, 1889, 205, 213, footnote); it is given as 255 4 (Lenz, Z. anal. Chem. 1889, 27, 572), 254 (Buttle, Henry and Trevan, 1c), 261 2 (Jungfleisch and Léger, Compt. rend. 1901, 132, Cinchona Micrantha. For purposes of quinine published values being [a]_b +225 in alrohol. manufacture it ranks as by product and rau bo The addition of chloroform, which enables a manuscture it ransa as a 57-product and ran be readily obtained from the cube mother-hoped stronger redution to be made, races the specific from which quinne sulphate has been crystallised, edition (+25%), \$1/40\$ achieved by taking solvantage of the spanning solvability of \$1.51.50 \, a how \$(-3)^1 + 263.75\$. Cuchonize by taking solvantage of the spanning solvability of \$1.51.50 \, a how \$(-3)^1 + 263.75\$. Cuchonize by taking solvantage of the spanning solvability of \$1.51.50 \, a how \$(-3)^1 + 263.75\$. ether and in alcohol. The cinchonidme is first is soluble in 3,810 parts of water at 10°, 3,670 at removed as tartrate, the mixed residual bases 20°, and 2,500 at 100°; it dissolves in 115 parts

chloroform (1), although its solubility in chloroform alone is only about one-third of that in alcohol. Cinchonine is fairly soluble in boiling amyl alcohol and in hot, but not cold, benzene. The addition of alkalis to aqueous solutions of cinchonine salts produces an amorphous pre-cipitate which gradually becomes crystalline, and which is stated to be more easily soluble in solvents than the crystalline form. Cinchonine begins to sublime at about 220° and boils with some decomposition above its melting-point; in a vacuum, however, it is volatilo unchanged. It is non-fluorescent and does not give the thalleioquin reaction. Unlike cinchonidine, it forms a soluble tartrate. Like quinine, cinchonine is bitter to the taste. Although cinchonine is a fairly strong dineidic base, aqueous solutions of its salts are liable to hydrolyse and to deposit the base. On oxidation with potassium permanganate it gives riso to cinchotenine, and with chromic acid it furnishes einchoninone, cinchoninie neid, meroquinene, eincholoiponie acid, and loiponic acid (for fuller descriptions of these products of oxidation, see below). On catalytic hydrogenation, hydroeinchonine (q.v.) is formed, while heating the acid sulphate with a little water at 140°, or boiling a solution of einchonine in dilute acetic acid, gives rise to cinchotoxine (cinchonicine) (sec below). With strong mineral acids various addition and transformation products are formed; and with amyl alcoholie potash cinchonine enters into dynamie equilibrium with *epic*inchonine, *ep*icinchonidine, and cinchonidine.

Cinchonine Sulphate, B₂·H₂SO₄,2H₂O, crystallises from water in the form of hard, transparent prisms, which become anhydrous at 100°, and then have m.p. 200° (decomp.). It is readily soluble in alcohol, less so in water (70 parts cold, 12 parts at 100°), and dissolves in chloroform to about the same extent as in water (distinction from quinine and cinchonidine). The salt dissolves in 2,300 parts of other at 25°. In 97% alcohol it shows [a]_b +193·3°-0·374c, where c=grams of base in 100 c.c. The optical relative in chloroform is the 12°.

rotation in chloroform is +133°.

Cinchonine Acid Sulphate, B.H₂SO₄,4H₂O, large, colourless prisms, is difficult to crystallise, being very soluble both in water and in alcohol.

Cinchonine Hydrochloride, B·HCl,2H₂O, crystallises from water in efflorescent prismatic crystals, which contain only 1H₂O when dried in the nir. The anhydrons salt melts at 217°–218° (decomp.) (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1919, 41, 819). It has $\{a\}_{0}^{10.5} + 177.4^{\circ}$ (ccal-083 anhydrous salt in water); $\{a]_{1}^{25} + 133.6^{\circ}$ (ccal-087 in chloroform); $\{a]_{1}^{15} + 165.5^{\circ}$ (2.425c in water] +214° (1.72c in water containing 2 equivalents of HCl) +172.7° (ccal in 97% alcohol), and +163.6° (ccal in 97% alcohol). The salt is soluble in the cold in 22 parts of water, in 1 part of alcohol, and 290 parts of ether. It also dissolves in enlaroform.

Among other salts of cinchonine may be mentioned the dihydrochloride, hydrobromide, in needles, has m.p. dihydrobromide, hydriodide, dihydriodide, $\pm 226.0^{\circ}$ (c=0.5 in hydriodate, nitrate, oxalate, d-tartrate, and chloride melts at 283°.

and in 18 parts of a mixture of alcohol (4) and bitartrate, l-bitartrate, benzoate, two citrates, chloroform (1), although its solubility in chloroform alone is only about one-third of that in alcohol. Cinchonine is fairly soluble in boiling salts are known.

Cinchonine Methiodide,

 $C_9H_6N-CH(OH)-C_7H_{11}N(CH_3I)-CH=CH_2$

is formed with a little difficulty from the components and melts at 254° (Claus and Müller, Ber. 1880, 13, 2290, footnote).

Cinchonine Ethiodide,

 C_9H_8N —CH(OH)— $C_7H_{11}N(C_2H_5I)$ — $CH = CH_2$,

is prepared by the gradual addition of ethyl iodide to a boiling solution of einchonino in alcohol in the dark. It crystallises from water in white, silky anhydrous needles, decomp. 260° (Claus and Kemperdick, Ber. 1880, 13, 2286).

Isomerio Cinchonine Ethiodide,

 $C_9H_6N(C_2H_5!)$ —CH(OH)— $C_7H_{11}N$ — $CH = CH_2$,

is made by heating 40 parts of einchonine hydriodide with 15 parts of alcohol and 15 parts of ethyl iodide for 2 hours under pressure. The product (base) crystallises from water in orange needles, and from alcohol in orange prisms, m.p. 184° (indef.) (Skraup and von Norwall, Monatsh. 1894, 15, 40). It is exidised with potassium permanganate to the ethiodide of einchoninic acid (Skraup, ibid. 433), and with a further molecule of ethyl iodide readily gives cinchonine diethiodide,

C₀H₆N(C₂H₅I)—CH(OH)—C₇H₁₁N(C₂H₅I)—CH=CH₂, also formed from the white monoethiodide, or from cinchonine, by heating at 150° with ethyl iodide in a scaled tube. It crystallises from water in large, dark yellow prisms with 1H₂O, decomp. 264°. It is readily soluble in water, and less soluble in alcohol than the white monoethiodide (Claus and Kemperdick, Ber. 1880, 13, 2928)

Acetyleinehonine,

 $C_9H_6N-CH(O-COCH_3)-C_7H_{11}N-CH=CH_2$,

results from the action of acetic anhydride at 60°-80° for 2 hours. Amorphous powder (Hesse, Annalen, 1880, 205, 321; Hilditch, J.C.S. 1911, 99, 238; Seekles, Rec. trav. chim. 1923, 42, 71).

Benzoylcinchonine,

 $C_{2}H_{4}N-CH(O-COC_{5}H_{5})-C_{7}H_{11}N-CH=CH_{2}$

is prepared by the action of excess of benzoyl chloride on einchonine. It crystallises in prisms from ether and melts at 106°-107° (Léger, Bull. Soc. chim. 1893, 9, 714; see also Skraup, Monatsh. 1895, 16, 163, and Seekles, Rec. trav. chim. 1923, 42, 71).

a-Chlorohydrocinchonine, better a-chlorodihydrocinchonine (component of "hydrochlorocinchonine"),

C₉H₆N—CH(OH)—C₇H₁₁N—CHCI—CH₃,

is isolated as the dihydrochloride from the products of the reaction of concentrated hydrochloric acid and cinchonine (Goodson, J.C.S. 1935, 1096). The base crystallises from alcohol in needles, has m.p. 236° (decomp.) and [a] $_{\rm D}^{20}$ +226·0° (c=0·5 in N-HCl). The dihydrochloride melts at 283°.

The mother-liquors from the crude dhydro-chloride contain a chlorohydroxine to make, which 1917, 50, 127); it can be distilled without is epimene with the a-derivative at the carbon decomposition. It crystallizes from ether in atom carrying the chlorine atom. It is purified is epimene with the a-derivative at the carbon atom carrying the chlorine atom. It is punfied in the form of its tartrate, which is crystallised from methyl alcohol. The base, which crystal lises from alrohol in minute needles, has mp 223° (decomp), and [a] +1760° (c=05 m N. HCl). The dibydrochloride melts at 276° (decomp.) and the tattrate, which crystallises from methyl alcohol with 2H O melts, when anhydrous, at 212º (decomp).

Bromohydrocinchonine, better bromodihydro cinchonine (hydrobromocinchonine),

first prepared from cinchoniao by Skraup (Annalen, 1890, 201, 324) and Comstock and Koenigs (Her. 1887, 20, 2520), is obtained from einchonine or a or \$ 100cinchonine, alloeinchonine, or a or \$ bydroxy by drocinchonine of a bydrog with 10 parts of 50% hidrohomie acid at 100° (Skraup, Copony and Medanich, Monatsh 1900, 21, 512, Zwerger, 1sed, 1903, 24, 119; Léger, Bull, Soc chim. 1918, [1r], 23, 133, 323; 1919, [1v], 25, 260) The base has m.p. 185°. the dihydrobromide er stallises from dilute bydrobromic acid in rolourless prisms, mp 2425-243° (Zwerger), [a], 1149 1° (c=1 989 in water) (Liger) The French author also mentions a (a) stereousomene bromobydro compound, +127 3°

a Iodohydroeinchonine (hydroiodoeinchonine), C,H,N--CH(OH)--C,H,,N--CHI--CH,,

is formed in predominating proportion when 18 formed in predominating proportion when cunchomno or a Meenchonino is bested at 100° with hydrodic acid (ppr. 17). The base, microscopic colourieae leakets, has mp. 1837–180° (Mecomp.). The hydrochloride has [a]_p + 180° (Mecc, tol. 1918, (pr. 23, 249) as 1800/Mecc, tol. 1918, (pp. 23, 249) as 1800/Mecc, tol. 1918, (pp. 23, 249) are 1800/Mecc, tol. 1918, (pp. 23, 249) are 1800/Mecc, tol. 1918, (pp. 23, 249).

at the sedme carbon atom, and is formed from allocinchonine and \$ isocinchonine. Its hydro-chloride has [a]o + 170.5° (Léget, le)

a 110Cinchonine (einehoniline),

is obtained by boiling einchonino with four times its weight of 50% sulphurse acid (Jung-hydrocinchonine by means of alcoholic potash

[a]17 +53 1° (c=1 m 97% alcohol), and +62 6° in water containing 4 equivalents of HCI. It is almost insoluble in water, but dissolves readily in alcohol, ether, acetone, benzeno or chloroform. It is a strong bitertiary base, which is alkalino even to phenolphthalein. Its oxygen atom has the ether function, for it cannot be acylated, and does not react with phenyl-hydrazine. The hydrochloride, B HCl,3H,O, has m p 226°, and [a]D +5° (c=1 of anhydrous salt in water). On heating its acid sulphate to 140°, a-1sociachonine undergoes the " Pasteur " reaction (Skraap and Zwerger, Monatsh, 1900, 21, 535; Ksas, sbid, 1905, 26, 119) giving rise to "a 150 of cinchonicine," which, in the light of the researches of Suszko and his collaborators on the assquandines, must be formulated as

(see Dubas, Konopnicki and Suszko, Rocz. Chem 1933, 13, 464). On boiling with dilute acetic scid, a secunchonine is converted into hydroxyhydrocinchotoxine,

(Rabe and Böttcher, l c.).
β 190Cinchonine (cunchonigine), a stereoisomendo of the preceding sabstance, is probably epimene with it at carbon atom "5" (see section 5d, p 149c) and is usually formed side by side with it, as well as from it by the action of sulphune acid (see references above, and Jungfleisch and Leger, Compt. rend. 1888, 106, 357; Ann. Chim. 1920, 14, 105 et seq.; Hesse, Annalen, 1893, 276, 88; Skraup, Monatsh. 1899, 20, 571; 1900, 21, 512). It is even more stable than a mounthonine. It exhibits dimorphism and separates from ether in transparent anhydrous crystals, belonging eithe to the orthorhombic or throthombic systems. The two forms are reversibly interconvertable. β isoCinchonine has mp. 130 7°, [a]17 -60 1° (c-1 in 97% alrohol), and -36° in water containing 4 equivalents of HCl. It is almost ansoluble in water, but readily soluble in most organic solvents including ether. It is a strong hitertisry base, but weaker than its a isomeride, being alkalino to himus, but not to phenelphthalein. It contains no hydroxyl or Leto group. The hydrochloride melts at 213°, and is soluble in about 50 parts of cold water; hydrounchonne by means of alcohole potash and the property of [a] -65 2° (c=1, anhydrous salt, water). On

acid at 100° (Skraup and Egerer, ibid. 1903, 24, epimeride it forms a diacetyl derivative, and is

allo Cinchonine (apocinchonine, pseudocinchonine).

$$C_3H_6N-CH(OH)-C_6H_{10}N>C=CH-CH_3$$

is an isomerido of the preceding two substances, and is usually formed along with them. It is a hitertiary base, forms acyl derivatives and adds the elements of hydrogen halide, giving rise to a small proportion of the same substances, which, under like conditions, are formed from cinchonine: the greater proportion of the resulting halogen derivatives are the same as these produced from β -isocinehonine. It erystallies from alcohol, has m.p. 216°-218° and [a]_D¹⁸ $\pm 150^{\circ}$ (c=1 in alcohol). It is characterised by the sparing solubility of the base in ether, and of the neutral sulphate in water (Jungfleisch and Léger, Ann. Chim. 1920, 14, 145 et seq., in which are further references to allocinehonine). Although a geometrical isomeride of alloeinchonine is unknown, a hase, m.p. 232°-234°, not identified with certainty, by Jungsleisch and leger (ibid. 181), probably constitutes missing substance.

a-llydroxyhydrocinchonine, better a-hydroxydihydrocinchonine,

$$C_9H_6N$$
— $CH(OH)$ — $C_7H_{11}N$ — $CH(OH)$ — CH_3 ,

is formed together with the preceding substances when cinchonino is heated with 50% sulphurie acid (Jungfleisch and Leger, Compt. rend. 1887, 105, 1255; 1888, 106, 68; 1889, 108, 952; Ann. Chim. 1920, 14, 71 et seq.; Léger, Compt. rend. 1918, 166, 903; 1919, 168, 404). It crystallises in colourless prisms, m.p. 252° (decomp.), $[a]_D^{18} + 182.6^{\circ}$ (c=1 in 97% alcohol). It is characterised by its sparing solubility in ether and ready solubility in 50% alcohol. It is almost insoluble in water, but dissolves readily in acetone, benzene or chloroform. a-Hydroxyhydrocinchonine gives rise to a diacetyl derivative. It is a strong bitertiary base alkaline to phenolphthalein. The neutral hydrochloride, B·HCl, H₂O, has m.p. 230° and $[\alpha]_D^{13} + 174.4°$ (r=1 in water with 1 equivalent of HCl). The salt is very sparingly soluble in water.

 β -Hydroxyhydrocinchonine, better β -hydroxydhydrocinchonine, is the epimeride of the foregoing substance at earbon atom " 5" (see section 5d, p. 149c). The two epimerides are formed together (see references above, and Jungfleisch and Liger, Compt. rend. 1894, 119, 1268; 1919, 168, 404; Ann. Chim. 1920, 14, 89 et seq.). It crystallises from 90% alcohol in small colourless, prismatic needles, m.p. 273° (decomp.) $[a]_{D}^{15} + 187.2^{\circ}$ (c=1 in 97% alcohol), or +203.1° in water containing 2 equivalents of HCl. Although it is insoluble in water, β -hydroxyhydrocinchonine is not immediately precipitated by alkalis from aqueous solutions of its salts, but separates in a fine mass of needles after some hours. It is readily soluble in alcohol, but not in acctone. It has a low solubility in chloroform, although this solvent will take up a good deal of the base at the moment of precipitation from its

alkaline to phenolphthalein. A number of salts have been described.

a- and β-Cinchonhydrine, components of "δ-einchonine," are substances of disputed constitution, the formulæ in question being C18H22ON2, C19H24ON2, and C19H22ON2. They are formed together with the einchonine isomerides described above, when hromohydroeinchonine base is heated with dilute alcohol, or with alcoholic potash (Jungfleisch and Léger, Compt. rend. 1894, 118, 29; Ann. Chim. 1920, 14, 158, 162, 165; Léger, Compt. rend. 1919, 169, 797; Cordier von Löwenhaupt, Monatsh. 1898, 19, 472; von Arlt, ibid. 1899, 20, 440; Langer, ibid. 1901, 22, 157). The a-isomeride erystallises from ether in prisms, m.p. 144.4°, $[a]_D^{10} + 195.8^{\circ}$ (c=1 in water containing 2 equivalents of HCI), or +139.8° in alcohol. The β -isomeride is similar. It crystallises from ether in prismatic needles, m.p. 155-8°, [a]_D²¹ $+106^{\circ}$ (c=1 in water containing 2 equivalents of HCl), or $+72\cdot2^{\circ}$ in alcohol. The einchonhydrines are said to ho the analogues of niquino (see under QUININE), and to be tertiarysecondary bases; they form diacetyl-derivatives.

A number of other "transformation products" of cinchonine are described in the literature. Most of these, if not all, can be ignored; somo are identical with, or are impure forms of, one or other of the substances described above; many are ohvious mixtures of one or more of the authentic compounds with one another, or with hydrocinchonine (derived from commercial cinchonine). For a valuable summary of the position in 1899, which is still largely valid owing to the fact that most of these spurious isomerides had been prepared before that time, see Skraup, Monatsh. 1899, 20, 571.

Cinchotoxine (cinchonicine),

$$C_9H_6N-CO-CH_2-CH_2-C_5H_9N-CH=CH_2$$
,

occurs in commercial quinoidine, the question of its pre-existence in the bark being in dispute. It is hest obtained artificially from einchonine or cinchonidine, either by melting (or heating with glycerol at 210°) the acid sulphate or tartrate, of either base, or, best perhaps, by the "wet" process of von Miller and Rohde (Ber. 1895, 28, 1064), in which einehonine is boiled with 14 parts of 7% acctic acid for 33 hours. Cinchotoxine crystallises from absolute ether at 0° in long, thin prisms, m.p. $58^{\circ}-59^{\circ}$, $[a]_{D}$ $+46.5^{\circ}$ in chloroform (c=2), +47.2° (Howard and Chick, r. infra), and +57.7° (Roques, v. infra), in alcohol; +35.6° (c=1 in water containing 4 equivalents of HCI). It is readily soluble in ether (2 parts), alcohol, benzene or chloroform, less so in petroleum; it dissolves in 564 parts of water at 18°, but is more soluble in aqueous solutions of ammonium salts, from which it expels ammonia; it is a strong base which absorbs carbon dioxide from the air. Cinchotoxine forms a crystalline hydriodide, oxalate, tartrate, nitrate (m.p. 160°), and double salts with zine, cadmium, and platinic chlorides. salts: the greater part of the alkaloid is soon Cinchotoxine is a tertiary secondary base, which deposited in the crystalline state. Like the alforms crystalline alkylhalides. It is a ketone,

forming a yellow phenylhydrazone, m.p. 145°, reduction in a suitable manner, einchoninono from a musture of alcohol and light petroleum [grees rise to cunchonne, epicinchonine, epi-cunchonine is non fluorescent and does not einchonidine, and einchondium. respond to the thallerooun reaction, but gives a magnificent purple coloration with diazobenzenesulphonic acid in presence of alkali. It reacts with amyl nitrite in presence of sodium ethoxide to form an isonitroso- derivative, and with pitrous acid gives a nitrosamme (N nitroso-derivative). (See Pasteur, Compt. rend 1853, 37, 110; Hesse, Annalen, 1868, 147, 241, 1873, 166, 277; 1875, 178, 253; Howard, J.C.S. 1872, 25, 101; Howard and Chick, J.S.C.I. 1909, 28, 55; Roques, Bull. Soc. chim. 1895, [10], 13, 1005; Ann. Chim 1897, 10, 234)

epiCinchonine, C13H22ON2, is epimerie with cinchonine about the earthinol (CHOH) group It is obtained by reducing einchonimme (100 g), dissolved in a solution of sodium ethoxide (Na 84 g in alcohol 2,280 ce), with almmining powder (84 g) The epicinehonine is isolated from the ether soluble fraction of the stereoisomerides formed, with the aid of tartane, dibenzoil, and diamsoyl. tartane acids. It erratallises from ether in long, colourless prisms, pointed at their ends, mp 82°-83°, [a]22 + 120-3° (c=0 8064 in alcohol). It is non fluorescent, and very soluble in all the usual organic solvents (Rabe and others, Annalen, 1932, 492, 253) Cinchoninone.

C.H.N-CO-C.H.,N-CH=CH., is obtained in very low yield by the gentle oxidation of einchonine or einchonidine with chromic acid (Rabe, Ber 1907, 40, 3655, 1009, 41, 62; Annalen, 1909, 364, 339) It is best prepared indirectly from einchotoxine The toxine (58 g.) in cold 5 hydrochloric acid (200 c c.), covered with a layer of other and well stirred, is first N brominated by introducing in a thin stream a cold solution of bromine (32 g.) in 6% sodium hydroxide solution (400 g.) (i.e. NaOBr) The N. bromotoxino (m p 153°) crystallises from ether, and is converted to emthoning to be treating a hot solution of le in 21 e c. of alcohol with 3 c c, of a cold sodium ethoxide solution containing Na 0-15 g (Rabe, Ber. 1911, 44, 2089) Cinchoninono erystallises from ether or from 50% alcohol in well defined, faintly yellow prisms, mp 126°-127°. It exhibits mutarotation, the final values being 0 +76 1° (c ≈ 3 302 m 99% alcohol), and [a]16 +74 7° (c=3 305 in benzene) (Rabe and others, Annalen, 1910, 373, 110) Cinchonmone 15 readily soluble in alrohol, ether, benzene or chloroform, but not in water, and sparingly soluble in light petroleum. It dissolves alightly with a yellow colour in aqueous alkalis (enolisa-tion). The hydrochloride (mp. 252°-253°) is very soluble in water. The initial values of the optical rotation of this salt depend on whether it has been prepared from a fresh or stale solution of the base; the final values, however, are the same : [a]14 +166 6° (c=1 656 m are the same : [a] +166 6° (c=1 656 in peroxide, and the ablehadic substances in chloroform) and +664° in water (Rabe and question. The latter are amorphous Cinchonothers, i.e.). Chehoninone forms an amorphous in all has $[a]_D + 100.5^\circ$ in chloroform; the oxime, and a benzoyl-derivative (derived from acctyl derivative has $[a]_D + \frac{43}{12}.2^\circ$, and the the enol form), which crystallises from petroleum | benzoyl derivative -72 4° (Seekles, Bec. trav. (b p over 80') in white needles, m p. 133'. On chim. 1923, 42, 69).

Cinchonine Chloride,

CaHeN-CHCI-CaHIIN-CH-CH

is prepared by treating dry cinchonine hydrochlorade (33 g) in chloroform (300 c.c.) with phosphorus pentaehloride (50 g) in the rold (Rabe and others, Annalen, 1910, 373, 101) The product (mp. 72°) crystallises from ether or from aqueous alcohol with 2H,O, which is lost over snlphuric acid; the anhydrous base then melts at 110°, and has [a]13 + 55 7° (c= 1 975 in 99% alcohol), and +62 20 (c=2 007 in chloroform). It is readily soluble in most organic solvents, but not in water (Koenigs, Ber. 1880, 13, 286). Its hydrochloride has [a] +49 5° (c=1.5555 in water).

Desoxycinchonine. C.H.N-CH,-C,H,,N-CH-CH,

is made by reducing emchanise chloride ilis-solved in 20 parts of 10% sulphuric soid, with 14 parts of iron filings at room temperature for 72 hours. The desoxy-hase er; stallies from light petroleum, has m p 91°, and [a], + 1793° (c=2 025 in 99% alcohol), and +194 3° in chloro form (Rabo and others, Annalen, 1910, 373, 103, 106).

Ciachene.

C.H.N-CH-C < C.H.N-CH-CH,

is formed when cinchonine or einchonidine chloride is boiled with alcoholic potash, It crystallises from light petroleum in colourless leaflets, m p. 123°-125°, and is destrorotatory. It can be distilled without decomposition (Kornigs, Ber 1881, 14, 1854; Koenigs and Comstock, ibid. 1885, 18, 1210; Koenigs, ibid. 1893, 28, 3146). apoCiachene, 4 (6 hydroxy 3 4 diethyl-

phenyl) quinoline,

 $C_H_N-C_H_1OH(C_H_1)$

(Kenner and Statham, J C.S 1935, 299), 11 obtained by beating cinchene with hydrobromic acid (sp gr. 1.49) at 180° under pressure for 6 to 8 hours (Locnies, Ber. 1881, 14, 1851; Kocnies and Comstock, 1814, 1883, 18, 1226). It crystallives from alrohol, and melts at 200 -210". baits and ethers of opocinchene are known (see Comstock and Koenigs, 161d. 1885, 18, 2379; 1887, 20, 2874; Koenigs, J. pr. Chem. 1900, [u], 61, 1). Cenchoninal,

 $C_{\epsilon}H_{\epsilon}N - CH(OH) - C_{7}H_{11}N - CH = O.$ and its act! (acetyl and benzoyl) derivatives, C.H.N-CH(O-COR)-C,H,1N-CH-O. are obtained from einchoning and from its acy derivatives by the action of ozone in cold chloroform solution. The resulting ozonides furnish on hydrolysis [formaldchyde], hydrogen

Cincholenine.

$C_0H_6N-CH(OH)-C_7H_{11}N-CO_2H_7$

is obtained by oxidising a cooled solution of cinchonine in dilute sulphurie acid with potassium permanganate. It crystallises from water with 3H₂O, m.p. 197°-198°, [a]_D¹⁵ +115.5° (c-2 of trihydrato in a mixture of 2 vols. chloroform and 1 vol. alcohol), +175.5° in dilute sulphuric acid (Hesse, Annalen, 1875, 176, 233; Skraup, ibid. 1879, 197, 379). Cinebotenine is soluble in hot water, but sparingly in alcohol. It dissolves readily in dilute acids and caustic alkalis, and is precipitated from the latter by carbon dioxide. As an acid, cincbotenine forms salts, esters and an acid chloride. Like einchonine it can be acylated, but does not react with the halogens or with their hydracids. It is stable to cold permanganate, but is rapidly oxidised by the hot reagent. With chromic acid the same oxidation products (but no meroquinone) result as in the like oxidation of cincbonine.

Cinchoninic Acid, quinoline-4-carboxylie acid. C₀H₀N—COOH, was first made by Caventou and Willm (Bull. See. chim. 1869, [ii], 12, 214) by the oxidation of einehonino with potassium permanganate. It is the most readily isolated of the exidation products of einchonine, cinchonidine, their hydro-derivatives, and many of the derivatives and degradation products of these. It can be prepared by the exidation of cinchonine with chromic acid (Koenigs, Bor. 1879, 12, 97) or nitrie acid (Claus and Muchall, ibid. 1885, 18, 362) or by oxidation with nitrie acid of the condensation product of lepidino with formaldchydo. It has been synthesised by tho interaction of isatin with acctaldoximo under tho ngency of 40% potassium hydroxide (Pfitzinger, J. pr. Chem. 1902, [ii], 68, 263; cf. Kaufmann, and Peyer, Ber. 1912, 45, 1805; Thielepape, ibid. 1938, 71, 387). Cinchoninic acid crystallises from water in needles with 1H2O, or in prisms with 2H₂O, depending on the conditions, and becomes anhydrous at 100°. It has m.p 253°-254°, is insoluble in ether, and sparingly soluble in water and in alcohol.

Lepidine, 4-methylquinoline, C,H6N-CH3, was first isolated from the products of dry dis-tillation of cinchonino with potash (Greville Williams, 1855), and has been synthesised by heating on the water-bath a mixture of equimolecular parts of aniline, acctone, and methylal (or formaldehyde) saturated with hydrogen chloride (C. Beyer, J. pr. Chem. 1886, [ü], 33, 418). Lepidine can be prepared by heating hydroeinchene or cinchene with 25% phosphoric acid at 170°–180° (Koenigs, Ber. 1894, 27, 1501; 1800, 23, 2077). Lepidine is a hygrogenia 1890, 23, 2677). Lepidine is a hygroscopic, sternutatory oil. h.p. 264.2°/760 mm., which folidifies on cooling and then melts at 9°-10°. It is sparingly soluble in water and is volatile in steam.

Oximinorinylquinuclidine, vinylquinuclidone-time. HO-N-C < C. H10N-CH=CH2, is formed together with cinchoninic acid from

cinchoninone (or with quininic acid from quininene), when these ketones, dissolved in cold alcoholic sodium ethoxide, are treated

of ether and light petroleum (b.p. 40°) in white needles, or from cthyl acctate in colourless prisms, m.p. 146°-147°. The oximo is readily soluble in water, alcohol and benzenc, fairly soluble in ether and ethyl acctate, and sparingly solublo in light petrolcum. It is volatilo in steam, and sublimes below its melting-point. On hydrolysis with hydrochloric acid, tho bieyelie system undergoes rupture, and there are formed bydroxylamine and meroquinene (Rabe, Ber. 1908, 41, 68; Annalen, 1909, 365, 361).

Meroquinene, 3-vinylpiperidine-4-acetic acid,

CH2=CH-C5H2N-CH2-CO2H,

is present among the products of oxidation of cinchonine, cinchotoxine, and quinine with chromic acid (Koenigs, Ber. 1894, 27, 1501; Annalen, 1906, 347, 196, 198). It is readily prepared by heating einchene or quinene with 25% phosphoric acid at 170°–180° (Koenigs, Ber. 1894, 27, 901, 904; Annalen, 1906, 347, 194, 195). Its N-methyl derivative is very readily obtained by subjecting isomitrace. readily obtained by subjecting isonitroso-Nmethylcinchotoxine to the Beckmann rearrangement, when the easily isolable. N-methylmeroquinene-nitrile is formed (Rabo and Ritter, Ber. 1905, 38, 2770). From a mixture of metbyl alcohol and ethyl acetato meroquineno separates in almost colourless crystals, m.p. 223°-224°, $[a]_{n}^{20} + 27.6^{\circ}$ (c=10 in water). It is very soluble in water, very sparingly so in cold alcohol, and almost insoluble in other or chloroform. Its hydrochloride has m.p 146°-148°, and the N-nitroso-derivative melts at 67°. On reduction with zine and hydriodic acid, meroquinene is converted to cincholoipon (see under Hydro-cinchonine, below) (Koenigs, Ber. 1902, 35, 1350), and on oxidation with cold potassium permanganate in acid solution it gives rise to Skraup's cincholoiponic acid. Mcroquinenc is an amphoteric substance; its ethyl ester is a strong monoacidic base, while its N-acctyl and -nitroso- derivatives are monohasic acids.

d-β-Cincholoiponie Acid, 3-carboxypiperidine-4-acetic acid, HO₂C-C₅H_BN-CH₂-CO₂H, results from the oxidation either with potassium permanganate or chromic acid or both, of all tho vinyl-alkaloids, of the toxines and tenines, of meroquinene, of eincholoipon, and of numerous meroquinene, of eincholoipon, and of numerous other transformation and degradation products of the cinchona alkaloids (Skraup, Monatsh. 1888, 9, 783; 1889, 10, 39; 1895, 16, 175; 1896, 17, 365; with Würstl, ibid. 1899, 10, 225; with Pum, ibid. 227; Schniderschitsch, ibid. 1889, 10, 54, 57; Würstl, ibid. 67, 70; Koenigs, Ber. 1895, 28, 3150; Annalen, 1906, 347, 208). It has been synthesised by Wohl, Losanitsch and Mang (Ber. 1907, 40, 4698; 1909, 42, 627). Cincholoiponic acid crystalliese 1909, 42, 627). Cincholoiponic acid crystallises from water in prisms containing 1H₂O (m.p. 126°-127°) which is lost at 120°-125°. Tho anhydrous acid melts at 225°-226°, and has $[a]_{0}^{20} + 30 \cdot 1^{\circ}$ (c=4 in water). It is very soluble in water and readily so in alcoholic hydrogen chloride, but it is insoluble in absolute alcohol and in other. Its hydrochloride has m.p. 192°with a little more than one molecular proportion of amyl nitrite. It crystallises from a mixture soluble in cold water and in alcohol, more

sparingly so in hydrochloric acid. Cincholoiponie By H,SO, 9H,O. It forms two methiodides: acid behaves as a monobasic acid, but its N- one crystallises with 3H2O and is white: nitroso- and N-acetyl derivatives manifest full dibasic properties. On oxidation with cold potassium permanganate solution it gives rise to losponic acid.

piperidine-3 4-dicarboxylic Losponse Acid, acid, C,H,N(CO,H), is obtained as described above (Skraup, Monatsh. 1896, 17, 376). It is also found in small amount in the mother houers from the cincheleipenic acid chtained by the ordation of cinchonine. Loiponic and crystal-hies from water in prama, mp 259-260. It is sparingly soluble in cold water, and dis solves in about 20 parts of hot water; it is almost insoluble in hot alcohol, readily soluble both in acids and in alkalis, and, on titration, behaves as a monobasic acid. It forms a nitroso-derivative, m.p. 167-168°, and its hydro-chloride (m.p. 216°-220°) has [a]_D about + 12° (c=833 in water) (Koenigs, Ber. 1897, 30,

1330). Cinchatine, See Hydrocinchonine, p. 162d. Cinchataxine. See un der Cinchonine, p. 157c. Conchairamidine, CarHas Os Na occura in the bark of Remijia Purdicana together with cinehonine, cinchonamine, concusconine, chair-amine, chairamidine, and conchairamine It was molated by Hesse (Annalen, 1884, 225, 256). For a scheme of separation the original paper should be consulted (15rd. 211, 218) It is obtained as a crystalline sulphate from the filtrate remaining after the chairamino has been separated as hydrochloride, and the conchair as thioeyanate, Conchairamidine separates from its solutions as an oil, and this gradually crystalleses with 1H1O. The anhydrous base has mp. 1140-1150, and [a]th -60° (c=3 in 97% alcohol). It is very soluble in alcohol, ether, chloroform, benzene, or acetone It is a weak base, neutral to litmus Concentrated sulphuric and gives a dark green colour. The hydrochloride, B HCl,3H,O, sulphate, B. H.SO, 14H.O, and other erystal-line salts have been described.

Conchairamine, C₂₂H₁₄O₄N₂, was solated by Ilesse (Annalen, 1884, 225, 246) from the raixture of alkaloids occurring in the bark of Remissa Purdicana After cemoving concusconing as sulphate, and chairamino as hydrochloride, the conchairamino is precipitated by potassium thio-cyanate. The base crystallises from alechel in thick, colourless prisms, B H2O C, H2OH, m p. 82°-66°. The alcohol of crystalisation is ex-pelled at 100°, leaving the monehydrate (m.p. 108°-110°) which, in its ture, becomes anhydrous and amorphous at 115°. The anhydrous form melts at about 120°, and has [a] 15 +68 4° (c=2 m 97% alcohol). An acetic acid solution of conchairamine, precipitated with ammonia, yields white eristalling flocks of the base, which alcohol. A solution in acetic acid gives a dark green coloration with concentrated nitrie acid, and sulphure acid produces a brown solution turning dark green. The base is neutral to trically grouped prisms containing 2H.O. htmus, but forms well-defined grafulation salts, which can only be expelled at 129-125. From such as the hydrochloride, B-HCl, and subplate, dilute alcohol it crystallises with 1/3H₂O.

the other, yellow, has 1H,O

Concusconine, C23H26O,N2, was isolated as the aulphate by Hesse (Annalen, 1884, 225, 234) from the alkaloids in the bark of Remina Purdicana. The base crystallises from alcohol in monoclinie needles containing 1H2O, mp. 144°; on further heating the melt solidates, and re melts at 206°-208°, [a] +40 8° (c=2 of the hydrate in 97% alcohol), but Howard and Chick (J.S C.I. 1909, 28, 55) record the value + 19 57°. Concuscommo is insoluble in water. very aparingly soluble in cold alcohol, and readily soluble in ether or chloroform. A solution in acctie or hydrochloric acid gives a dark green colour with concentrated nitric seid, With concentrated sulphurie acid a blush green colour is produced, turning to olive green on warming Concusconino is a weak base, neutral to litmus, and its salta are for the most part amorphous. It gives riso to two methodides, and contains two methoxyl groups,

Conquinamine, C10H21O, N, 15 present together with quinamine in small quantity in most emchons barks, and was first isolated by liese (Ber. 1877, 10, 2158). It is found in the alcoholic mother liquors from which quinamico has been crystallised, and is isolated by means of light petroleum, and purified by erystallisation of the ratrate, oxalate, or hydrobromide. It crystallises from alcohol in long prisms or pyramids, m p. 123°, [a] + 2014° (c=1 in alcohol). It is fairly soluble in alcohol, ether, or benzene, but not in 50% alcohol or in water, It resembles quinamine in behaviour, and seems to be closely related to it, for on melting its exalate, or on heating with hydrochloric and (sp gr. I-125) quinamione and apoquinamine are formed, which are also phtained from quinaomic. A number of crystallino salts of conquinamine have been described (Hesse, Annalen, 1881, 209, 62; Oudemans, told, 38),

Conquinine, See Quinidine (p. 1896). Cuprelne, C19H22O2N2, occurs together with other cinchona alkaloids (but not einchonidine) in the se called " Cupres bark" (Remijia pedunculata), which is no longer collected commercially, and there being no known usy of preparing it artificially from other alkaloids, cupreino is now unobtainable. It was first isolated by Paul and Cownley (l'harm. J. 1881, 15, 221, 401) from "homoquinine," a losse compound of quante and cupcerne, at first mistaken for a new alkaloid cesembling quinine;

homogunane is claimed to have been discovered by Hesse's assistant, Tod (see Hesse, Ber. 1882, 15, 857; 1883, 16, 60), but it was first described in the published literature by l'aul and The name Cownley (Pharm. J. 1881, 12, 497). " ultraquinine" was proposed for it by Whillin contain 1 H₂O, and delay drate and melt at 110°, (that, 497) and "homoquinino" by Howard and It is soluble in chloroform, ether, and hot Hodgkin (that, 528; J.C.S. 1882, 41, 66); see also Hesso (Annalen, 1884, 225, 95; 1884, 226, 240; 1835, 230, 55).

Cupreme crystallises from other in concen-

-175.5° (c=1.2 in alcohol). For rotations in various alkaline solutions, sec Oudemans (Rec. trav. chim. 1890, 9, 171). It is readily soluble in alcohol, but sparingly so in ether, ehloroform, or benzene. As a phenol, enpreine dissolves freely in aqueous caustie alkalis, from which it is reprecipitated by earhon dioxide, but is not reachly soluble in ammonia. Cupreine is nonfluorescent, but gives the thalleioquin reaction, and a red-brown coloration with ferrie chloride solution. Stereochemically, cupreino is einchonidine with a hydroxyl group in the 6-position of the quinoline nucleus, or the phenol of which quinine is the methyl ether, for eupreine gives rise to quinme when heated with 11 molecules of methyl nitrate in presence of sodium methoxide in a sealed tube at 100° (Grimaux and Arnaud, Compt. rend. 1891, 112, 774). With methyl iodide there are formed cupreino methiodide, dimethiodide, quinino methiodide or dimethiodide, depending on the conditions. With hydrochloric acid it gives rise, like quinine, to "apoquinine" (see under Quinme) (Hesse), and on entalytic hydrogenation to hydroeupreine (see under Hydroquinine) (Gieinsa and Halberkann, Ber. 1918, 51, 1325). It forms a diacetyl derivative.

The salts of eupreine are colourless, and the acid salts afford colourless solutions, but the neutral salts dissolve in water with an intenso yellow colour. The sulphate, B2. H2SO4, crystallives from water in anhydrous needles (Howard and Chick, J.S.C.I. 1909, 28, 55), m.p. 257° (decomp.) (Giemsa and Halberkann, l.c.), soluble in 813 parts of water at 17°, and in 209 parts at 100°. It is also sparingly soluble in alcohol. The acid sulphate, B.H2SO4, contains 1H2O according to Hesse, and Howard and Chick, but 2H2O according to Oudemans (Rec. trav. chim. 1889, 8, 147) and Giemsa and Halberkann. It separates in large, transparent, faintly yellow, prismatic plates, $[a]_D^{22}$ -197.9° (c= 1.1821 in water), and is soluble in about 70 parts of cold water. A tetrasulphate is also known. Cupreinc hydrochloride, B.HCl, H2O (colourless needles), dissolves in about 50 parts of cold water, and has [a]_D¹⁷ -154.8° to -157.1° in water (c=0.57 to 0.86), and -167.3° to -169.7° in alcohol (c=0.9 to 1.42). For other salts the papers of Hesse and of Ondemans already cited should be consulted, and for physiological action of cupreine see Grimaux and Laborde (Compt. rend. Soc. Biol. 1892, 44, 605).

The procedure whereby Grimaux and Arnaud (l.c.) succeeded in methylating cupremo to quinino enabled them also to prepare the following series of homologues of quinme:

Quirethyline, C₂₁H₂₆O₂N₂, ethylcupreine, (Compt. rend. 1891, 112, 1364), m.p. 160° (dry), [a]_D -1694°. The sulphate, B₂·H₂SO₄, H₂O₅ dissolves in 397 parts of water at 15°. According to Giemsa and Halberkann (Ber. 1918, 51, 1332) the base has m.p. 165°-166°, and [a]_D²⁰ -158.9° ia alcohol.

Quinpropyline, C.,H.,O.N., n-propyl-

The anhydrous base melts at 198° and has [a], 1892, 114, 672), m.p. 164° (dry). The sulphate, B₂·H₂SO₄,1½H₂O, is soluble in 454 parts of water at 13°.

Quinisopropyline, C₂₂H₂₈O₂N₂, isopropyleupreine (ibid.), m.p. 151°; B₂·H₂SO₄,H₂O, dissolves in 367 parts of water at 10°.

Quinisoamyline, $C_{24}H_{32}O_2N_2$, isoamyleupreino (ibid.), is amorphous, m.p. 167°. $B_2\cdot H_2SO_4$ is soluble in 4,170 parts of water at 11.5°.

Unlike cupreine, the above homologous ethers fluoresee in dilute sulphurie acid solution. They are said to be more powerfully antipyretie, but more toxic than quinine (Grimaux, Laborde and Bourn, Compt. rend. 1894, 118, 1303).

Cuscamldine, of undetermined composition, occurs in Cusco bark (Cinchona Pelletierana). It is amorphous, and is obtained together with cuscamine (which it closely resembles) by tho addition of nitrie acid to an acetic acid solution of the total alkaloids of the bark. The two bases are separated from one another as oxalates. The salts of cuscamidine are amorphous and are more soluble than those of cuscamine (Hesse, Anualen, 1880, 200, 304).

Cuscamine, of undetermined composition, is obtained from Cusco bark (sec under Cuscamidine, obote). I terystallises from alcohol in prisms, m.p. 218°, is readily soluble in ether and in hot alcohol, but sparingly so in cold alcohol. It gives a yellow colour with concentrated sulphuric acid, turning brown on warming; on adding molybdic acid a bluish-green colour is obtained, turning brown on heating, and violetbrown on re-cooling. With nitrie acid a yellow coloration is obtained. Cuscamine is not a strong base, but many salts are known, some of which are crystalline (Hesse).

Cusconidine, of undetermined composition, is an amorphous alkaloid occurring in Cusco bark (Cinchona Pelleticrana). It is obtained from eusconine mother-liquors (see below). salts are amorphous (Hesse, Annalen, 1880, 200. 303).

Cusconlne, C23H26O4N2, occurs in Cusco bark and is obtained as sulphate after removing the arieine (q.v.) as acctate. It was discovered by Leverkoehn in 1829. It erystallises from ether in white leaflets with 2H₂O. The water is lost at 100° and the dry base has m.p. 110°, $[a]_D^{15}$ -54.3° (c=2 in 97% alcohol). It dissolves in 35 parts of ether at 15°, more easily in alcohol or acctone, very readily in eliloroform, and is insoluble in water. Cusconino gives the samo colour reactions with sulphuric acid and with ammonium molybdate as does aricine (q.v.). It is a weak base: its salts have an acid reaction, and are amorphous (Hesse, Annalen, 1877, 185,

Diclochonine (dicinehonicine), CasH44O2N4, has the same empirical composition as emchonine, and although its molecular weight is half that indicated by the formula, Hesso prefers the double formula by analogy with diconquinine. It occurs in Cinchona rosulenta and in C. succirubra together with the other principal cinchona alkaloids. Being soluble and amorcupreine (Grimaux and Arnaud, Compt. rend. | phous it tends to pass into the so-called " quino-

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dine" fraction in quinine manufacture, and is alcohol, ether and light petroleum in anhydrous, recovered therefrom as the thiocyanate. It is purified by crystallisation of the hydrochloride. The free base is a yellow amorphous substance, m p 40°, [a] +91.7° (c=1 52 in 97% alcohul). It is very soluble in alcohol, ether, acetone, chloroform, or benzene, but not in alkalis. It does not give the thalleroquin reaction. It is a atcong base, and its hydrochloride, B 2HCL crystallises well (Hesse, Annalen, 1885, 227, 153) On heating with hydrochlorie seid, it is converted into the isomene, amorphous diapocinchonine, soluble in ether, alcohol, or ehlocoform, [a]u +20° in alcohol (Hesse, Annalen, 1880, 205, 333]. Diapocinchonine is a mixture of isomerides containing a- and \$ woeinchonine (Jungfleisch and Leger, Compt. rend. 1892, 114, 1192; Hesse, Annalen, 1893, 276, 118).

Diconquinine (diquinicine), CasH45O2N2, the essential constituent of commercial "quinoidine" (q r.), is prepared by removing from quinoiding the toxines in the form of exalates A chloroform extract of the residue, a red-brown amorphous mass, constitutes "diconpunnine" (llesse, Ber 1877, 10, 2155) It is dextrorotatory, fluorescent in dilute sulphuric acid, and shows the thalleroquin reaction. Its salts are amorphous. According to Hesse diconquinine is to be considered as derived from the condensation of two molecules of conquinine (quinidine) with the climination of one molecule of water. There seems to be little evidence for this view, and Hesse himself found that the molecular weight of diconquinine corre-

sponded to the angle (C₁₀) formula Hydrocinchonidine (cuchamidine, dihydro-cuchonidine), C₁₁H₁₂ON_p, was first isolated by Font and Böhringer (Ber. 1881, 14, 1270) by oxidising commercial cinehonidine, which contains hydrocinehonidine, from Cinchona Ledgersana, with cold potassium permanganate in acid solution. The cinchonidine being converted to the amphoteric substance einchoteni-dine (see under Cinchonidine), and the hydroemchonidine remaining for the most part unchanged, it could be readily isolated. Hadro canchonidine is best prepared by the establish hydrogenation of canchonidine, with which it is identical except that it contains an ethyl sidechain in place of the vinyl group of the latter. It is identical with Hesse's cinchamidiue (Ber. 1881, 14, 1683.) which he isolated from the mother liquors of emchandine sulphate.

lly decementation erystallises from hot dilute alcohol in hexagonal leaflets, mp 229"-230", [a] -98 4' (c=2 m 97% alcohol); -144 6° (c=0 735, s.e. an M/40 solution in 0-1N-H,SO4) (Buttle, Henry and Trevan, Brochem J. 1934, 28, 438). It is almost insoluble in water, very sparingly soluble in ether and ehlocoform, and more sparingly soluble in alcohol than is cinchonidine. It is non fluorescent and does not show the thallesoquin reaction.

The hydrochloride, BHCl,2H,O (Hesse, Annalen, 1882, 214, 1) has [a]_D -804° (c= 1-19 of hydrated salt in water). It is very soluble in water or alcohol. According to lleidelberger and Jacobs (J. Amer. Chem. Soc. 1919,

microscopic needles and plates, m p. 2020-2030 [a] -89-4° (c=1-197 in water), and is readily soluble in chloroform, and fairly soluble m acctone.

The sulphate, B. H.SD., 7H.O. needles, is soluble in 57 parts of cold water.

Hydrocinchonidine is stable, it is not readily oridised by potassium permanganate, nor isomerised by the action of hot mineral acids, but on melting the acid sulphate, hydro-cmehotoxinn (see under hydrocinchonine) is formed and no heating with hydrochloric acid in scaled tubes epihydeoeinehonidinn is formed, On oxidation with cheomie acid it gives rise to emehonmic acid.

epiHydrocinehonidine, C13H24ON2, is epi-merie with hydrocinehonidine about the carbinol (CH-OH) group, and is obtained either by the eatalytic hydrogenation of epicinchonidine, or by the catalytic reduction of hydrocinchoninnan or by inversion of hydrocinchonine (q r) with amyl alcoholie potassium hydroxide (Rabe, and others, Annalen, 1932, 492, 254, 255, 263). It crystallises from other in colonrless needles, m p. 106°, [a]20 +483° (c=1-004 in 99% alcohol) It is very readily soluble in ether and in accione, and more so than epshydrocinchonme. epillydrocinchonidine was first described, but not named, by Vereinigte Chininfabriken, Zummer & Co, G m.b H., in O.P. 330813, 1920. Hydrocinchonidine Chloride,

CaH,N-CHCI-C,H11N-C,H1 was prepared by Cohen and King (Proc. Roy. Soc. 1938, B, 125, 54). The hydrochlonds crystallises from alcohol, mp. 233°-225° (decomp.), [a]20° +24° (c=0 5 in water).

Dezoryhydrocinchonidine,

C.H.N-CH,-C.H.IN-C.H,

has been made by the catalytic hydrogenation of cinchonidine chlorida (Cohen and King, Proc. Roy. Soc. 1938, B. 125, 56). It has also been obtained from epshydrocinchonidine chloride by reduction with iron and sulphune acid (Rabe and others, Annalen, 1932, 492, 256), and from the hydrazone of hydrocinchoninone by heating at 180° with potassium hydroxide (ibid. 263, 264, 265). In the last preparation a mixture nf desoxyhjdrocinehonidine and desoxyhydroemchoume (see under Hydrocinchonine) is formed, the desoxyhydroeinehonidine being seolated as the danasoyltartrate from actions.
The base has m.p. 52° (bbd. 256) or 42° -44° (bbd. 256), and [e]₀ -21·2° (e-0 4 or 0 8 in alcohol).
The hydrochloride, from a mixture of methyl alcohol and ether, has m p. 186°-188°, and [a]

+9-0° (c-1-08 in water). Hydrocinchonine (einchotine, dibydrocin-ehonme), C₁₉H₂₁ON₂, was first isolated from Remiya Pardicona by Caventou and Willia (Annaken, 1870, Suppl. 7, 247). It occurs in most cinchona barks, and is a common component of commercial curchonine, whenro Herse obtained it by oxidation with potassium permanganate in sulphurio acid solution at 0 (Annalen, 1873, 168, 256). He therefore re-41. 821) the salt crystallises from a mixture of garded it as an oxidation product of einchoning "cinchonine," and that the effect of the reaction with permanganate is to oxidise the latter alkaloid, not to hydrocinehonine but to the amphotoric substance einchotenine, and thus to facilitate the isolation of the hydrocinchonine which is left largely unattacked (Annalen, 1879, 197, 362). Other methods for isolating hydrocinchonine depend on the crystallisation of the hydriodides (Hesse, Annalen, 1890, 260, 213, 220; 1898, 300, 45; Pum, Monatsh. 1895, 16, 70), on heating with dilute sulphurie acid (the cinchonine becomes greatly altered, whereas tho hydrocinchonine, remaining unchanged, can be readily separated from the resulting mixture) (Juagileisch and Leger, Bull. Soc. ehim. 1888, 49, 747: 1901, 25, 881), on fractional crystallisation of the dihydrobromides (Buttle, Henry and Trevan, Biochem. J. 1934, 28, 437), and best, on the formation of a compound between einchouine, but not hydrocinehonine, and mercurie acetate (Thron and Dirscherl, Annalen, 1935, 515, 252). Hydrocinchonine is most veniently prepared, however, by the catalytie hydrogenation of cinchonine, from which it differs only in containing an ethyl side-chain in place of the vinyl group of the latter.

Hydrocinchonine crystallises from alcohol in prisas and scales. Its melting-point is variously given between 267° (Buttle, Henry and Trevan, l.c.) and 278° (Jungsleisch and Léger, l.c.). Hesse found 268°-269°, and Skraup 277·3°; [a]_n in alcohol, between +189-8° (Rabo and others, Annalen, 1910, 373, 100) and +204-5° (Hesse, ibid. 1898, 300, 46); [a]_n in aqueous acid solution between +225-8° (Buttle, et al.) and +227.7° (Jungfleisch and Leger). Hydrocinchonine dissolves in 1,360 parts of cold water, but is more soluble in boiling water. It dissolves in 534 parts of ether, or 220 parts of alcohol at 20°, and is much less soluble than einchonine

in an alcohol-chloroform mixture.

Hydrocinchonine is stable towards mineral acids, and is unable to react additively with hydriodic acid, but heated in a scaled tubo with 25% hydrochloric acid, it gives rise to some epihydrocinchonino (eec below); it can be sulphonated by means of concentrated sulphurie acid. Boiling with diluto acctie acid affords hydrocinchotoxine (see below). It is not readily attacked by potassium permanganate, but with chronic acid it is oxidised to hydrocinchoninono (see below), einchoninic neid, eincheloipon (see below), cincholoiponie and loiponie acids (see under Cinchonine).

Hydrocinchonine Hydrochloride, B.HC1,2H2O (Forst and Böhringer, Ber. 1881, 14, 437; von Arlt, Monatsh. 1899, 20, 431; Heidelberger and Jacobs, J. Amer. Chem. Soc. 1919, 41, 826) crystallises from water in needles, m.p. 216° (von Arlt), or 221°-223° (dry) (Heidelberger and Jacobs), $[a]_0^{25} + 159.3°$ (c-0.741 in water). It is soluble in 47 parts of cold water.

The sulphate, B. H.SO4, crystallises in prisms, leaflets, or needles with 2, 6, 9, but ordinarily with 11 or 12H2O, depending on the conditions. The dihydrate has [a] +164.4°

but Skraup showed later that it pre-exists in the soluble in about 30 parts of cold water and in "cinchonine," and that the effect of the reaction about 10 of boiling water. The dodecahydrate is efflorescent. Many other salts of hydroeinchonine are known such as the neutral and acid hydrobromide and hydriodide, acid hydrochloride and sulphate, two tartrates, and a thiocyanate.

Hydrocinchotoxine (dihydrocinchotoxine, hydrocinchonicine, cinchotinetoxine, cinchoticine),

$$C_9H_6N-CO-CH_2-CH_2-C_5H_9N-C_2H_5$$

is obtained by boiling hydrocinchonine with diluto acetie acid (Kaufmann and Huber, Ber. 1913, 46, 2919; Kaufmann, Rothlin, and Brunnschweiter, ibid. 1916, 49, 2303). It has also been obtained by "partial synthesis" (Rabe and Kindler, ibid. 1918, 51, 1360) from einehoninic ester and benzoylhomocincholoipon ethyl ester; (for the subsequent synthesis of homocincholoipon, see Rabe and others, ibid. 1931, 64, 2493). It is a viscous oil, $[a]_D^{23} + 8.8^{\circ}$ (c=6.17 in alcohol). Its N-benzoyl-derivative has m.p. 124°, and the dipicrate of its phenylhydrazone melts at 215°. The free toxine, which is readily soluble in ether or in dilute alcohol, can be brominated, and is then on treatment with alkali converted into hydrocinchoninone (see below). Hydrocinehotoxine gives purple colorations with nitrobenzeno containing nitrothiopliene and with diazobenzenesulphonic acid.

epiHydrocinchonine, C₁₀H₂₄ON₂, is epimerie with hydrocinchonine about the carbinol (CH·OH) group. It was first obtained but not named by Vereinigte Chininfabriken, Zimmer & Co., G.m.b.H., G.P. 330813/1920, by the reduction of hydrocinchoninone with aluminium or zine in alcoholic solution in presence of sodium ethoxide. It can also be obtained from hydrocinchoninone by catalytic reduction, from hydrocinclionine by epimerisation with amyl alcoholic potash, and from cpicinchonine by catulytic hydrogenation (Rabo and others, Annalen, 1932, 492, 252-255, 263). In all cases except the last, a mixture of hydrocinchonine, hydrocinchonidine, epihydrocinchonine, and epihydrocinchonidine is produced. The first two naturally occurring alkaloids are readily removed by taking advantage of their sparing solubility in ether. The two residual epi-bases are separated mechanically and by fractional crystallisation from ether and from acctone. cpiHydrocinchonine crystallises from acctone in colourless, tapering prisms, m.p. 126° , $[a]_{D}^{19}$ +88.4° (c -1.0012 in 99% alcohol). It is readily soluble in other or acctone, and very readily so in alcohol or benzene.

Hydrocinchoninone,

C_pH₆N—CO—C₇H₁₁N—C₂H₅,

is obtained in low yield by the gentle oxidation of hydrocinchonino with chronic acid (Rabe, Naumann and Kuliga, Annalen, 1909, 364, 349), but is best prepared indirectly from hydrocinchotoxine by way of the C-bromo derivative (Kaufmann and Huber, Ber. 1913, 46, 2920; Kaufmann and Haensler, ibid. 1917, 50, 703; Rabe and Kindler, ibid. 1918, 51, 1365). (c=25 in alcohol). The anhydrous salt, which It separates in pale yellow crystals from 60% absorbs 2H₂O on exposure, has m.p. 195°, is alcohol, has m.p. 138° (Rabe) or 130°

(Kaufmann) and [a]n (final value) on the average | einehene). It is isomeric with, but distinct from +76° in alcohol (c=from 2 to 4). The hydrochloride, which is quite white, crystallises from alcohol and ether and has m.p. 265°. The oxime is amorphous. For the mutarotation of hydrocinchoninone, see Rabe and others (Annalen, 1910, 373, 118).

Hydrocinchonine Chloride,

is made by treating dry hydrocinchonine hydrochloride in chloroform with phosphorus pentachloride (Koenigs and Hoerlin, Ber. 1894, 27, 2291; Jacobs and Heidelberger, J. Amer. Chem. Soc. 1922, 44, 1087) Crystallised from etherit has ro p. 850-870 (indef) or with \$ 5H.O from aqueous acetone, m.p. 70° (undef). The latter form, on drying in a desiceator, becomes amorphous and has [a]14 +36-4" (c-1-001 m alcohol). It is very readily soluble in organic colvents. The hydrochloride, m p. 227°-228°, +48 8° (c=1 107 in water), ervstallises from 85% alcohol, and is aparingly adjuble in cold water or alcohol, but more readily so in tho boiling solvent. The hydrobromide is very eraringly soluble in water. Desoryhydroeinekonine,

is made either by reducing hydrocinchonine chloride with iron in dilute sulphuric seid (E. G. W Schmidt, Dissertation, Jens, 1913, Jaroha and Heidelberger, J Amer. Chem Soc 1922, 44, 1088), or by hydrogenating desoxycanchomne or emchene using a palladium-calcium carbonate catalyst (Schopf and Schmidt, Annalen, 1928, 465, 127), or by heating the hydrazone of hydrocinchoninone with powdered potassium hydroxide (Rabe and von Riegen, Annalen, 1932, 492, 263). Desoxyhydrocinchonine base crystallises from moist ether or from aqueous acetons with 2H,O, mp. 60°, [a] +113 8° (c=1-014 in alcohol). The water is lost in dry air or in a desiceator, and the base becomes amorphous. It separates in the anhydrous condition from light petroleum or from absolute ether, and then has m.p. 74", [a]" +147° in alcohol. It is easily soluble in organic solvents. The hydrochloride crystallises from alcohol in narrow glistening plates and prisms, rop 197°-199°, [a]_p²⁴ +69·3° (c=1 008 in water). The salt is fairly soluble in alcohol, very readily so in methyl alcohol or in chloroform, but less so in acctone. The hydrodule has m p. 230°-231°. Hydrocinchene,

$$C_1H_1N-CH-C < C_1H_{16}N-C_1H_{5}$$

is prepared by boiling hydrocurchonine chloride with alcoholic potash (Koenigs and Hoerlin, Ber. 1894, 27, 2291; Koenigs, ibid. 1895, 28, 3146 ; J. pr. Chem. 1900, [11], 61, 44 ; see also Koenigs, Ber. 1894, 27, 1504). It crystallises an ethyl side-chain in place of a vinyl gro 'P from aqueous methyl alcohol in leaflets, mp It has been aynthesised by Rabe, Huntenbur. from aqueous methyl alcohol in leasiets, mp It has been aynthesised by Rabe, Hante 145'. With 25 % phosphoric acid at 170' it is Schultze and Volger (Ber. 1931, 64, 2487). hydrolysed to lepidine (q r. under Cinchonine) and emcholospon (see below), but does not react plates) or alcohol (needles), and is said to contain with hydrobromic acid (distinction from 25H₂O which is easily expelled, but the exist-

desoxycinchonine.

Oximinochylquinuclidine, ethylquinuchdoneoxime, HO-N=C < C1H10N-C1H1, is formed together with einchonine acid from hydroconchaninone, when this ketone, dissolved in cold alcoholic sedium ethoxide, is treated with a little more than one molecular proportion of amyl netrete (Rabe and Naumann, Annalen, 1909, 365, 363). It crystallises in white needles from a mixture of light petroleum (h p. 60°-65°) and ethyl acetate, and has m.p. 133°-134°. It is readily soluble in water, alcohol, ether, or ethyl acetate, but less so in light petroleum. On hydrolysis with hydrochloric acid, the bicycle system undergoes rupture, and there are formed hydroxylamine and cincholorpon.

Cincholorpon, 3-ethylpiperidine-4 acetic acid, C.H., C.H., N-CH., -CO,H., was first discovered by Skraup (Monatas 1888, 9, 603) the products of oxidation of cunchonine and quinine with chromic acid. It was shown by Koenies, however (Ber, 1894, 27, 1504; with Hoerlin, stid 2290), that the cancholospon was derived not from these alkaloids but from the hydro alkaloids present in the commercial materials. Cincholoipon can be prepared either by the hydrolysis of hydrocinchene with 25 phosphoric acid at 170°, or by exidation of hydrocanchonina with chromic acid (Koenigs, and Koenigs and Hoerlin, ic) It has also been prepared by the reduction of meroquinene (q v. ander Cinchonine) either with sinc and hydrodic and (Koenigs, ibid. 1902, 25, 1350), or better catalytically (Kaufinann, Zeller and Huber, shid. 1913, 48, 1830). It can also be prepared from hydroeinchoninone, by the action of amyl mitrite followed by hydrolysis (see Oximmoethylquinuclidine, abore). Its N. methyl derivative is readily obtained by audiceting monitroso-N-methyleinchotoxine to the Beckmann transformation; the easily isolable N-methyl derivative of the nitrile of cincholospon is then formed (Rabe and Ackermann, Ber. 1907, 40, 2013). Cincholospon crystallises from methyl alcohol and has m p. 236°. It is soluble in water. The hydrochloride consists of yellow rhomhic crystals, m p. 200°, readily soluble in water or in alcohol, but not in ether. The salt has [a] -5 6" (c=10 in water). Cincholoipon ethyl cater hydrochloride has mp. 158° and [a]22 +5.71° (c=8 29 in water).

Hydroquinidine, C, H, O, N, occurs in commercial quandine, cometimes in large amount. It was first isolated from this source by Forst and Böhringer by oxidation with potassium permanganate in acid solution at 0° (Ber. 1881, 14, 1934; 1882, 15, 520, 1656), the effect being to oxidise the unsaturated quinidine to the alkan-soluble quiterndine, and thus to facilitata the isolation of the hydroquinidine. It is best prepared by catalytic hydrogenation of quinsdine, from which it differs only in having

Hydroquindine crystallises from ether (stout

ence of such a hydrate is doubtful (Giemsa and Halberkann, Ber. 1921, 54, [B], 1201). The dry base melts at 169° and has $[a]_0^{20} + 231\cdot3°$ (c= 2.0216 in alcohol), and +299° (c=0.815, i.e. an M/40 solution in 0·1N- H_2SO_4). It is sparingly soluble in ether, but dissolves readily in hot alcohol. It gives the thalleioquin reaction, and fluoresces in dilute sulphuric acid solution. Under the influence of hot mineral acids it is demethylated to hydroeupreidine (v. infra), but does not undergo isomerisation. It is fairly stable towards potassium permanganate, but with chromic acid is oxidised to quinine acid (q.v. under Quinine), and eincholoipon (v. supra).

Hydroquinidine Hydrochloride, B·HCl, is readily soluble in water, melts at 273°-274°, and has $[a]_D^{26} + 183 \cdot 9^\circ$ (c=1·3 in water) (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1919,

41, 826).

Hydroquinidine Sulphate, B₂·H₂SO₄, separates from water above 20° in needles containing 3H₂O (Giemsa and Halberkann) or 2H₂O (Ilesse). The anhydrous salt, n.p. 214°, does not absorb water on re-exposure. When crystallised below 10°, the sulphate separates in prisms, and these are so efflorescent that it is difficult to determine their initial water content, which is given as 8 and 12H₂O, but may be as high as 18 H₂O. The hydrate stable in the air appears to be B₂·H₂SO₄.2H₂O, for the anhydrous salt reabsorbs that amount on exposure. Ilydroquinidine sulphate dissolves in 92·3 parts of water at 16°.

The dihydrochloride has been used to separate quinidine and hydroquinidine (Buttle, Henry and Trevan, Biochem. J. 1934, 28, 434); the dihydrobromide has been described by Emde (Ilelv. Chim. Acta. 1932, 15, 575). The neutral tartrate, B₂·C₄H₆O₆, 2H₂O, is readily soluble in water, but the acid tartrate, B·C₄H₆O₆, 3H₂O,

is sparingly soluble.

epiHydroquinidine, C₂₀H₂₀O₂N₂, is epimeric with hydroquinidine about the carbinol (CH·OH) group. It is obtained either by heating hydroquinine with amyl alcoholic potassium hydroxide, or by the eatalytic reduction of hydroquininone, or by catalytic hydrogenation of epiquinidine. In each case except the last, a mixture of hydroquinine, hydroquinidine, epihydroquinine and epihydroquinidine is produced. After removing the first two naturally occurring alkaloids as neutral and acid tartrate respectively, the two ϵpi -bases are separated as neutral dibenzoyltartrates from benzene. The ϵpi hydroquinidine so obtained from the more soluble component is finally purified by recrystallisation of the base from ether. epiHydroquinidine has been synthesised (Rabe and others, Ber. 1931, 64, 2499). It forms colourless, that prisms from ether, m.p. $1^{\frac{99}{2}}$, $[a]_{11}^{16} + 73.7^{\circ}$ (c:-0.3321 in alcohol), is very readily soluble in alcohol, fairly soluble in other, and is strongly fluorescent in sulphurie acid solution. Its dibenzoyltartrate, m.p. 1569-157', is sparingly soluble in water, other, benzene, or cold alcohol, but readily so in hot alcohol (Rale and others, Annalen, 1932, 492, 253, 256, 257, 261).

Hydroquinidine Chloride,

 $CH_2O \cdot C_9H_5N - CHC! - C_7H_{11}N - C_2H_5$

is prepared from dry hydroquinidine hydrochloride and phosphorus pentachloride in chloroform solution (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1920, 42, 1497). It crystallises from light petroleum in cream-tinted plates, m.p. $93\cdot5^\circ-95^\circ$, $[a]_D^{\circ 2}+20^\circ$ ($c=0\cdot6$ in alcohol). The hydrochloride, needles from alcohol, m.p. $208^\circ-209^\circ$, $[a]_D^{\circ 2}+39\cdot7^\circ$ ($c=0\cdot795$ in water), is readily soluble in water and in methyl alcohol, less so in ethyl alcohol or in ebloroform, and sparingly so in acctone.

Desoxyhydroquinidine,

 $CH_3O-C_9H_5N-CH_2-C_7H_{11}N-C_2H_5$

is obtained by reducing hydroquinidine chlorido with iron filings in dilute sulpburic acid (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1920, 42, 1497). It has also been made similarly from epihydroquinidine chlorido (Rabe and others, Annalen, 1932, 492, 258), and the following particulars are taken from these two sources (cf. Giemsa and Halberkann, Ber. 1921, 54, 1203). Desoxyhydroquinidine crystallises from 60% alcohol with 2H₂O, m.p. 85°-87°, [a]_p^{22.5} +167·3° (c=1·124 in alcohol). The anhydrous base melts lower at 68·5°-70°.

It is readily soluble in alcohol, acctono, chloroform, or benzene and less so in ether. It gives the thalleioquin reaction and fluoresees in dilute sulphurie acid solution (purple). Its

hydrobromide melts at 151°-152°.

Hydrocupreidine, C10 H24O2N2, was first obtained by Heidelberger and Jacobs (J. Amer. Chem. Soc. 1919, 41, 827) by the demethylation of hydroquinidine with 4 parts of boiling 40% hydrobronic acid at 125°. It can also be prepared from hydroquinidine with strong sulphuric acid, the resulting hydrocupreidine sulphonic acid being saponified by heating with dilute hydrochlorie or sulphurie neid (Giemsa and Bonath, Ber. 1925, 58, [B], 93), or directly by boiling with 60% sulphuric acid (Henry and Solomon, J.C.S. 1934, 1929) Hydrocupreidine crystallises from aqueous alcohol (Heidelberger and Jacobs) or from acctone (Henry and Solomon) as an indefinito hydrato which is difficult to dry without discoloration; hydrate melts at 170°. The anhydrous base melts indefinitely at 195°. According to Giemsa and Bonath the base crystallises from absoluto alcohol, and then has m.p. 193°. The specific rotations given for the dry base in alcohol are widely different: +253.4° (Heidelberger and Jacobs), -1.242.5° (Giemsa and Bonath), +227.2° (Henry and Solomon). The base is readily soluble in cold alcohol, less readily in chloroform or acctone, and sparingly in ether. It is non-fluorescent, and does not give a coloration with bromine water and ammonia. The hydrochloride crystallises from 50% alcohol in rosettes of needles which are not anhydrous, as stated by Henry and Solomon (l.c.), but contain 1H.O; the anhydrous salt melts at 232, and has [a]n +194° in water. The dilydrochloride sparingly soluble in dilute hydrochloric acid. For other salts and for the ethyl ether of hydrocupresdure, see Heidelberger and Jacobs (I c.), janhydrous salt melts at 200°-208°, and has [a]o and for other homologous alkyl ethers, see Ghosh and Chatterjee (J. Indian Chem. Soc. 1931, 8, 237; 1932, 9, 83; Buttle, Henry, Solomon, Trevan and Gibbs, Biochem. J. 1933, 32, 52).

Hydroquinine, C20H21O2N2, accompanies quinino in cinchona barks and in commercial quinine sulphate. It was first isolated by Hesse (Ber. 1882, 15, 856; Annalen, 1887, 241, 257, 286) from quinine sulphate mother-biguous, by taking advantage of the fact that hydroqumme acid sulphate, BH,SO4, or hydroquinine dihydrohromide, B2HBr, is more soluble in water than the corresponding quinine salt. Quinine can in this way be completely freed from hydroquinine, but the latter can only be prepared in the pure state by oxidising with potassium permanganate the mixture of quinine and hydroquinine obtained in the mother liquors; the quinine becomes oxidised to the alkalı soluble quitenine, and the hydroquinine, being left largely unattacked, esn then be readily isolated. It is, however, best prepared by the catalytic hydrogenation of quinine, from which it differs only in that it has an ethyl sidethain in place of the vinyl group of the latter This process has in the last decade or two acquired much technical importance, partly because of the demand for hydroquinine itself as a therapeutic agent, and partly because it con-atitutes the starting material for the manufacture of the hydrocupreme others (" optochin, "vuzin") Early examples of such catalytic processes can be found in the following patents: G P 234137/1911, 252138/1912, 267306/1913; U.P 3949/1012. For details of proresses, see CATALYSIS IN INDUSTRIAL CHEMISTRY, Vol. II, \ p 422

Hydroquinine hase is precipitated by sodium hydrovide solution from aqueous solutions of its salts, when it separates in amorphous flakes which gradually become crystalline and contain 2H.O. This form turns to a soft rubbery mass below 100°, but the anhydrous form, which crystallises from ether or benzene, melts at 17.2 3° (Lenz, Z anal Chem 1888, 27, 561) and has [a]_D -142 2° (c=2 4 in 95% alcohol) (Hesse), or -233 7º (c=0 815, se. an M/40 solution in 01N-H,SO, (Buttle, Henry and Trevan, Biochem. J. 1931, 28, 436) It is readdy soluble in most organic solvents, including ether, but not in water or alkalis; it nevertheless dissolves appreciably in ammonia solution Hydroquinine is fluorescent in ddute sulphuric acid solution, and gives the thallesoquin reaction It is relatively stable towards permanganate, but is oxidised by chromic acid to quininic acid (q v. under Quinine) and eincholomon (q v under Hydrocinchenine). On melting its acid aul phate, or on boiling with dilute acetic acid, it is transformed to hydroquinotoxine (e. infro) With strong mineral acids (HCI, HBr, HI, H.SO.) or aluminium chloride it nadergoes demethylation (see "Hydrocupreme" below), but not isomerisation. Ilydroquinine is latter to the taste, and is equal or superior in antimalarial activity to quinine.

-I23 9° (c=1-113 in water). It also crystallises from a mixture of aretone and other with 0 5H.O. It is readily soluble in water, alcohol or arctone, but not in ether (Hesse, Annalen, 1887, 241, 264 : Heidelberger and Jacobs, J. Amer. Chem.

Soc 1919, 41, 820) Hydroquinine Sulphate, B, H, SO, crystallises from water in needles containing 6H,O when stable, but probably with 8H,O at first. salt is soluble in 348 parts of cold water, and when anhydrous re-absorbs 4 or 6H,O on exposure. The acid sulphate, B.H.SO. crystallises with 3H.O and is extremely soluble in water. The dinydrobromide erystallises with 3 H2O, and the anhydrona salt has [a] 15 -152 5° in water. Many other salts of hydrogumine are known. It forms loose compounds with several other emehona alkaloids and also with anethole; it forms an amorphous aretyl- derivative, crystalline benzoyl- derivative (mp. 102°-107°), and a methiodide.

Hedrogumotoxine (hydrogumicine).

is prepared from hydroquinine either by heating the acid sulphate at 140° (Hesse, Annalen, 1887 244, 273) or by boiling in dilute aretic and (Rabe, Ber 1912, 45, 2927, Kaufmann and Huber, sold, 1913, 46, 2921; Heidelberger and Jacobs, J. Amer Chem. Soc. 1922, 44, 1092) It has been synthesised by Rabe and his coworkers (Ber. 1931, 64, 2497; see also Rabo and Kindler, 181d. 1919, 52, 1842). It is a yellow varnish, [a]D -17° (c=3 in water with 3HCl). easily soluble in alcohol, ether, or chloroform. Ita solution in dilute sulphurio a cid is yellow, but non fluorescent, and it gives the thillconum reaction The sulphate, B, H, SO,, crystalizes from a mixture of aqueous alcohol and acetone with SH₂O; the anhydrous salt has m p. 174°-176°, [a] = -83° (c=0 968 water) Oxalic acid precipitates an amorphous exclute, which is soluble in chloroform (distinction from quinotoxine); the dibencoyltarirate

melts at 170°, epiHydroquinine, CapHgaO1N2, in epimeric with by droquinine about the carbinol (CH OH) group. It can be prepared either from alcoholic potassium hydroxide, or from hydroquinone by catalytic reduction, or by the catalytic hydrogenation of epiquinine. In the first two cases there is formed a mixture of hydroquinine, hydroquinidine, epihydroquinine and epibydroquinidine. The first two naturally occurring alkaloids are removed as neutral and acid tartrate respectively; the two epi bases are then separated as neutral dibenzoyltartrates from benzene. The epihydroquinine so obtained from the more sparingly soluble component is finally porified by recrystallisation of its hydrochloride from acetone. Attempts to crystallise the base have so far fauled; it is a viscous oil, [a] +32° (r=0 95 in alcohol), readily soluble in organic solvents, and showing a strong blue Hydroquinine Hydrockloride, B HCl, 2H, O. fluorescence in sulpliuric acid solution. The crystallists from water in prisms. The dibenzo thattrate, mp. 151° crystallises from acctone, and is sparingly soluble in water, benzene, light petroleum or ether readily so in alcohol or chloroform (Rabe and co-workers, Annalen, 1932, 492, 253, 256, 257, 260).

Hydroquininone,

was first prepared by Irschiek (Dissertation, Jenn, 1913) by the gentle exidation of hydroquinine with ehromic acid. It is best prepared indirectly from hydroquinotoxine. By brominating in hydrobromic acid solution the Cbromotoxine is obtained, which, on treatment with sodium othoxido in alcoholic solution, gives rise to the desired ketone (Kaufmann and Huber, Ber. 1913, 46, 2922). It has been synthesised by Rabo and his co-workers (Ber. 1931, 64, 2497; see also Rabe and Kindler, ibid. 1919, 52, 1845). Hydroquininono crystallises from other in yellowish needles or leaslets, m.p. 99°; it shows mutarotation: [a]_D (final value) +73·15° (c=1·784 in alcohol) (Kaufmann and Huber; Rabo and Kindler; Irschick's ketone, prepared by direct oxidation, and Rabe's synthetic ketono havo lower rotations, +65.0° and +68.8° respectively). The pierate melts at 2240.

Hydroquinine Chloride,

is prepared by the action of phosphorus pentachloride on dry hydroquinine dihydrochloride dissolved in cold chloroform (Heidelberger and Jacobs, J. Amer. Chom. Soc. 1920, 42, 1491; Gieman and Halberkann, Ber. 1921, 54, [B], 1101). It crystallises from dilute alcohol or from petroleum spirit (b.p. 80°-90°) in creameoloured rhembs, m.p. 144°, $[a]_D^{21.5} + 42 \cdot 1^\circ$ (c= 1.365 in alcohol) or $\pm 43^{\circ}$ (c=3.0136 in alcohol). Hydroquining chloride is readily soluble in alcohol, acetone, chloroform, or benzene, less readily in ether, fairly soluble in boiling but very sparingly soluble in cold petroleum spirit. It is a light-sensitivo substanco which gives the thalleioquin reaction. Aqueous acid solutions are yellow, but non-fluorescent. The hydrochloride, B.HCl, m.p. 232°-233°, $[a]_D^{20.5} - 2.9^{\circ}$ (c=1.197 in water) crystallises from 50% alcohol, and is somewhat sparingly soluble in water or alcohol.

De soxyhydroquinine,

$$CH_{3}O-C_{9}H_{5}N-CH_{2}-C_{7}H_{11}N-C_{2}H_{5}$$

is obtained by reducing hydroquinine chloride with iron in dilute sulphuric acid solution (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1920, 42, 1492; Gienia and Halberkann, Ber. 1921, 54, 1198), or in the same manner from epihydroquinine chlorido Annalch, 1932, 492, (Rabe and 258). Deso: others, Desoxyhydroquinine crystallises from aqueous acctone in colourless, silky needles which contain 3H2O and melt at 69°, solidify on further heating, and melt again at 95°. The 3H.O is readily lost in a desicentor; the anhydrous base is oily, absorbs 2H2O from the air, and in doing so crystallises. The trihydrate has [a]_p²⁴ -77.5°

organic solvents; it gives the thalleioquin and fluorescence tests. The hydrochloride, B-HCl, erystallises from acetone in anhydrous needles, m.p. 179°-180°, $[a]_D^{225}$ -6.8° (c=1.034 in water). The salt dissolves readily in water, alcohol, methyl alcohol, ehloroform, or hot acctono or benzene, but sparingly in cold acctone or benzene.

Hydroquinene,

$$CH_2O-C_2H_5N-CH=C < C_6H_{10}N-C_2H_5$$

is prepared by boiling hydroquinino chloride with alcobolic potassium hydroxide (Giemsa and Halberkann, Ber. 1921, 54, [B], 1194). It crystallises from a mixture of ether and light petroleum in colourless rosettes of coffin-shaped needles. and on recrystallisation from dilute alcohol or acetone, separates with 2H2O. The anhydrous substanco melts at 78°-79°, and has [a]_n²⁰ $+29.5^{\circ}$ (c=2.678 in alcohol). It is readily solublo in the usual organic solvents, except in light petrolcum. Hydroquineno gives thalleioquin test, fluoresces in dilute nitrie acid solution, and is instantly attacked by potassium permanganate. The hydrochloride melts at 209°.

Hydrocupreine, C19H24O2N2, is prepared by demethylating hydroquinine with hot hydro-chloric (Hesse, Annalen, 1887, 241, 280), hydrobromie (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1919, 41, 821), or hydriodic acid (Pum, Monatsh. 1895, 16, 73), with aluminium chloride in nitrobenzene solution (Oberlin, Arch. Pharm. 1927, 265, 270), or with concentrated sulphurie acid, when a sulphonic acid is formed which is hydrolysed by hoiling with dilute hydrochloric or sulphuric acid (Gicmsa and Bonath, Ber. 1925, 58, [B], 89, 92). It can also be obtained by the catalytic hydrogenation of eupreino (Giemsa and Halberkann, ibid. 1018, 51, 1328), hut best, perhaps, by boiling hydroquinine with 60% sulphuric acid. Apart from demethylation, no change takes place, for hydro-quinine is regenerated when hydroeupreine is methylated by means of diazomethane (Oberlin.) Hydroeupreine is precipitated by means of

ammonia as a micro-crystalline powder containing 2H2O, but erystallises (if pure) from alcohol, dilute alcohol, or from a mixture of ehloroform and benzene in the anhydrous condition. From benzene alone it separates as a jelly. It becomes amorphous on heating below its melting-point, and this accounts for a divergence among the published figures; Hesse and Pum give m.p. 170°; Giemsa and Halberkann record 204°, while Giemsa and Bonath, seven years later, record 186°; according to Heidelberger and Jacobs hydrocupreine becomes a glassy mass filled with bubbles of gas at 185°-190°, and this does not become completely liquid until 230°. The specific rotation of hydrocupreine in alcohol is given variously between the extremes -148.7° (Heidelberger and Jacobs) and -159.2° (Giemsa and Bonath). Hydrocupreine is readily soluble in alcohol or chloroform, and as a phenol dissolves freely in caustic alkalis. It is sparingly soluble in ether or ethyl acetate, almost insoluble in light petroleum or water, and dissolves with difficulty in ammonia solution. (c-1-244 in alcohol) and is readily soluble in eupreine is non-fluorescent and does not give to have obtained the green colour by using -136 2° (c=1 005 in alcohol) bleaching powder. Aqueous solutions of the neutral salts, like those of cupreme, are of an isolated by Hesse (Ber. 1877, 10, 2162) from the intensely yellow colour, which is immediately hark of Cinchona Calisaya var. jaranica. It discharged on the addition of mineral acids, erystallises from water in rhombic leaflets, is Hydrocupreine hydrochloride (B HCI) eratal- readily soluble in other, forms a crystalline lises from water acidified with hydrochlone oxalate, and dissolves in dilute sulphuric acid acid in anhydrous needles, decomp. 255°-280°, melting at the latter temperature, [a]22 -1323° (c=0945 in alcohol), it is aparingly soluble in water, acetone, or chloroform, readily discolves in alcohol and very readily in methyl alcohol. The dihydrochloride, B 2HCLHO, melte at 221° The neutral sulphate, B, H, SO. is anhydrous and very sparingly soluble in water or in sleohol. The end sulphate, BH₃SO₃, separates with 4H₂O₄, the anhydrous salt, mp 20(3°, reabsorbs 1H₂O₄O₄, exposure The tarteste, B₃C₃H₄O₄,2H₂O₅, melts at 242°. Other salts prepared include the diliydre bromide and seid nitrate.

Hydrocupreine gives nee to a toxine, bydrocupremotoxine (hydrocupreicine), in the ordinary way, but a number of derivatives such as hydrocupreme bromide and desovabadro cupreme have been obtained by demethylating the corresponding hydroquinine derivatives. As a phenol, hydrocupreme, in common with cupreine, couples with diazonium solutions, giving rise to an interesting series of dyes.

A considerable amount of attention has been given to the alkyl others of hydrocupreme and hydrocupremotoxine since the discovery of their atriking bactericidal ection by Morgenroth and collaborators, who investigated the complete series from ethyl to cetyl (Cis) Those interested in the results of these biological investigations and their possible therapeutic application should consult. The Therajeutic Agenta of the Quinoline Group." by you Octtingen (Chemical Catalog to Inc. New York, 1931). "Handbuch der Chemotherapie." by Fischl and Schloss berger (Frielier's Medizinische Buchhandlung, Lengug, 1932) and 'Recent Advances in Chemotherapy 'by Findlay (J and A Churchili, London, 1930) 1 or the preparation of some of these aubstances, see Vereinigte Chimnfabriken, Zimmer & Co. G.m b H , G P 254712, 1912, 344140 (1921 . Girman and Halbertann, Ber 1918, 51, 1332, Heidelberger and Jacobs, J Amer Chem Soc 1919, 41, 824, 1922, 44, 1007, Slotts and Behnisch, Ber 1933, 66, [B], 360, and for the entire series up to C₁₁, see Buttle, Henry, Solomon, Trevan and Gibles, Biochem. J. 1938, 32, 51. Among the fatter known products are isomythydroenpreme dibydrochloride ("eucupin"), eer-octylhydro-cupreme dibydrochloride ("yuzin") and ethyl hwirocupreine hydrochloride (" optochin ").

which is recognised in the "I nited States Pharmacopera," M. The salt readily ersatal lises from a mixture of actions and ether, m p. 25."-254", [a]" -123 6" (c=0.959 in water). It

the thalleroquin reaction although Hesse claims bydrous base has mp. 123"-128" and [ali.

Javanine, of undetermined composition, was with an intensely yellow colour.

Niquidine - See under Quinidine.

Paricine. C15H15ON, was discovered by Winekler (Jahresbericht, 1845, 27, 338) in the barks of Buena hexandra and Cinchona lutes, Later, Hesse found it in C. succirubra of Darjeeling (Pharm J. 1870, 1, 344; Annalen, 1873, 166, 263) It is an amorphous, yellow powder, m p 116°, soluble in water, alcohol, other or light petroleum Its salts are also amorphous, and it s nitrate is insoluble Quanamine, C10H21O2N2, was isolated from the bark of Cinchona succirula by Hesse (Ber

1872, 5, 265, see also ibid, 1877, 10, 2157; Annalen, 1873, 166, 266, 1881, 207, 288). It occurs only in small quantities, but is widely distributed, one of the richest sources being Cinchona Ledgeriana Oudemans (Annales, 1679, 197, 49) found that the total alkaloids of C succirubra contained 45% of quinamine. It is best isolated from quinine sulphate mother liquors (see foregoing references and da Vrn. Pharm J 1874, 4, 609, Howard, and 1875, 5, 1) by removing quinine and cinchondine as tartrates, and curchonune as thiocianate The quinamine crystallises from a solution of the residual bases in 80% alrohol, or may be purified by crystallisation of the nitrate. The base crystallises in long, anhydrous needles, mp. 1720, [a]15 +93 4° (c=2 in chloroform), cr +104 5° in alcohol It is almost insoluble in mater, sparingly so in alcohol, and fairly soluble in ether, it gives a rich jellow coloration with concentrated aulphoric acid in the presente of a little mitrie acid, on diluting with water, tha colour changes through purple to pink. Quinamine is a monoacidic base, and its acid salts are amorphous; the neutral salts, of which a fairly large number have been prepared, crystallice well. It contains no methoral group, is readily oxideed, and is very unstable towards acide a variety of products being formed (fless Annalen, 1881, 207, 292 et eeq). Very mild treatment, such as boiling with 10 sulphune acid, or even standing at the ordinary temperature in 13% hydrochloric acid, soffices to isomen-e quinamine to quinamidine (m p 93' (alm +45" in alcohol). This is best prepared (seed 299) by heating quinamine with 5 parts of to tartaric acid solution for 2 hours To gether with quinamidine is always formed in smaller amount another isomeride, quinquicise (mp 109°, [a]p +38° m alcohol), which is obtained also from conquinamine (q t.), but in greatest amount when an alcoholic solution of quanamine seid sulphate containing a little plycerol as exaporated at 60°-80°, and the resulte beated at 100° for several minutes (idem, ibid, 303) is readily soluble in water or alcohol. The base. If the residue is heated at 1.31 -130' instead of on the market is also amorphous, but has been at 1011, an amorphous autotance predoquinamiobtained crystallino from toluene; the an- cine, C17H20O2N2, is formed (ideia, ibid. 303)

1.125) or with 25% sulphuric acid for 3 minutes, quinamino is dehydrated to apoquinamine (C19H22ON2, idem, ibid. 294) (m.p. 114°; base optically inactive, hydrochloride.

B·HCI, HO,

lævorotatory). apoQuinamine is formed in the same way also from quinamicine or conquinamino (q.v.). Quinamine and apoquinamine give rise to one and the same acetyl-derivative, which on hydrolysis furnishes the latter base. On heating at 140° with hydrochlorie acid saturated at -17° quinamine is converted into a rubbery mass, which is insoluble in dilute acids, or in

ordinary organie solvents.

Quinetum.-This name was used by de Vrij to describe the total alkaloids of C. succirubra, which in the early days of einchona-planting he recommended as the most economical method of utilising cinchona alkaloids. The preparation was also known as "cinchona febrifuge." When Cinchona succirubra was supplanted by species yielding bark richer in quinine, it became difficult to supply "quinetum" and the demand was met in two different ways. The residues of the total alkaloids of "factory bark," after the removal of quinine, were made into a quinetum substitute by the return of some quinine and the product was known as "einchona febrifuge." the product was known as emerion tenting the life variability in composition has frequently been commented upon (see, for example, Leaguo of Nations, Health Scetion, C.H. Malaria/158, Geneva 14/4/1931; Howard, Trans. Roy. Soc. Trop. Med. Hyg. 1925, 18, 358; Geodson and Henry, Quart. J. Pharm. 1930, 3, 220 and it is being like the state of the life of t 238) and it is desirable that it should be replaced by a standardised preparation such as totaquina (p. 132).

Cinchona febrifuge was sold principally to indigent natives in tropical countries. There was also a certain demand for quinctum in Europe, and under the new conditions this demand was met by mixtures of the crystalline alkaloids of cinchona, but as there was no standard this product was also variable (League of Nations, Health Section, I.c., 15-181, and the Malaria Commission of the League of Nations proposed in 1931 that the name quinctum should be restricted to a mixture in equal parts of quinine, cinchonine, and cinchonidine. standard is recognised in the "British Pharma-ceutical Codex," 1934, which requires that quinctum shall contain not less than 60% of quinino and cinchonidine as determined by the "British Pharmacopæia" method for totaquina (see p.132). The loss in drying is not to exceed 5%.

and the ash must be 1% or less.

Quinicine.—See Quinotoxine, under Quinine.

Quinidine (conquinine), C20H21O2N2, is a naturally occurring dextrorotatory stereoisotucride of quinine, and is stereochemically identical with cinchonine, except that it contains a methoxyl group in the 6-position of the quincline nucleus. It is found in small quantity in most cinchona barks, but especially in C. pitayensis, C. amygdalifolia, and C. Calisaya. Quinidine is now extracted on a commercial

When boiled with hydrochloric acid (sp.gr., cipitated together with the amorphous alkaloids in the "tails" of quinine manufacture, and so passed into the agglomeration of alkaloids termed by Sertürner "quinoidine" (1828). It was from this source that Henry and Delondro (1833) prepared an alkaloid which they termed quinidine," while Van Heyningen (1849) also isolated a crystalline alkaloid, isomerie with quinine, but called it " β -quinine," for Winekler (1847) had in the meantime used the name "quinidine" for another alkaloid. Pasteur (1853) renamed Winckler's alkaloid "cinchonidine" (the name which it now bears, q.v.), and reverted to Henry and Delondre's original name for the alkaloid here in question, which he isolated also from commercial cinchonine, and which de Vrij (1857) showed to be identical with Van Heyningen's β -quinine. To avoid confusion with Winckler's alkaloid (cinchonidine), Hesso (1865) proposed for the quinidine, now of unambignous identity, the name "conquinine," a name much used in Germany but not elsewhere. Tho alkaloid is now almost everywhere known as quinidine.

Quinidine is prepared from quinine sulphate mother-liquors, from which, on adding eaustie soda, it passes together with einchonidine into ether. The einchonidine is removed as tartrate, and the quinidine isolated as hydriodide. It is the least soluble of the cinchona hydriodides. The base is finally recovered, and recrystallised from alcohol. Quinidine so prepared contains large amounts (e.g. 30%) of hydroquinidine which can be removed either by Thron and Dirscherl's mereurie acetate process (Annalen, 1935, 515. 252), or by fractional recrystallisation of the dihydrochloride from water (Buttle, Henry and

Trevan, Biochem. J. 1934, 28, 434).

According to Hesse (Ber. 1877, 10, 2154; cf. Mylius, ibid. 1886, 19, 1773) quinidine crystallises from dilute alcohol in prisms containing 2.5H2O (losing 0.5H2O on exposure); from absolute alcohol it separates with 1EtOH; from ordinary ether in rhombohedra containing 2H2O; from absolute other with 3Et2O; from boiling water in leaslets with 1.5H,O; and from benzene it crystallises in anhydrous needles, m.p. 172°. The quinidine of the "British Pharmacentical Codex," 1934, 872, contains 2H₂O and melts at about 168°. The published rotations of quinidine are: +262° (c=2 in alcohol) (Hille, Arch. Pharm. 1903, 241, 77); +256° (c=3·2 in alcohol) (Ondemans, Annalco, 1870, 182, 45); +243·5° (c=0·77 in alcohol) (Rabe, ibid. 1910, 373, 100); +230° (c=1·8 in chloroform) (Hesse, ibid. 1875, 176, 224) and $+334.2^{\circ}$ (c=0.81, i.e. an M/40 solution in 0.1 N. H2SO4) (Buttle et al. l.c.). For other rotations, see Ondemans (I.c.) and Lenz (Z. anal. Chem. 1888, 27, 571). Quinidine is not very readily soluble in any one organic solvent; it dissolves best in alcohol, but is soluble also in other, chloroform, benzeno or cthyl acetate: it is almost insoluble in cold water, but dissolves in 750 parts of boiling water. It gives the thalleioguin reaction, and exhibits a blue fluorescence in dilute sulphurie acid. Quinidine closely resembles quinine both in chemical and pharmacological properties; both yield the reale, but at one time it was said to be pre- samo toxine, quinone, quininic acid, and

meroquinene; on oxidation with potassium] permanganate it gives rise to quitenidme (see below) and on catalytic bydrogenation to hydroquinidine (q r.). On heating with mineral acids it undergoes various changes, see- products of the reaction of concentrated hydroand opoquinidae and their cogeners being chloric and and quindine (Goodson, J.C.S. formed feet below). Ouinidine is not greatly 1935, 1097). The base crystallises in needless inferior to quinine in antimalarial efficacy. It has been recommended for use in place of the lavorotatory quinine in cases of "quinine idiosyncrasy

Quinidine Hydrochloride, B HCl, crystallises in asbestos like prasma containing 1H,O (Hesse, Annalen, 1868, 146, 362; 1875, 176, 225), [e]h +205 8° (4 928e in water), +212° (2 562c in 97%, alrohol). The anhydrous salt melta at 288-259°. It is readily soluble in alrohol or bot water, but only in 625 parts of water at 10°. It dissolves also in chloroform, but is almost

insoluble in ether.

Quinidine Sulphate, B. H.SO, crystallises from hot water in prisms with 2H,O (Hesse, Annalen, 1868, 146, 364; 1875, 176, 226; 1876, 182, 141). It is described in "British Pharma-copera," 1932, 364, as colourless, needle-like crystals. It has [a]n +179 5° (c=1 m water), +211 5° (c=2 m alcohol), or +184 2° (c=3 of anhydrous salt in chloroform). It is soluble in about 100 parts of cold water, 7 parts of boiling water, 10 parts of alcohol, or 20 of chlocoform It is almost insoluble in ether. The anhydrous salt reabsorbs 2H.O on exposuce and is light sensitiva The said sulphate is readily soluble in water

Quinidine Hydrodide, BHI, is precipitated as a crystalline powder when a solution of a quinidine salt is treated with potassium iodide. It is used for the isolation and estimation of quandine, being very sparingly soluble in cold water (1 in about 1,250) and also in alcohol or boiling water.

The neutral tartrate, B, C,H,O,H,O, dissolves in about 40 parts of water, the good tartrate B C.H.O., 3H.O (mp 100°) in 400 parts. Quinidine Methiodide,

CH,O C,H,N CH(OH) C,H,1N(CH,F) CH CH,,

is formed from the components in the cold, or more readily on boiling in alcoholic solution in the dark. It crystallises in colourless needles with 1 H₂O, and is readily soluble in alcohol or boiling water, but sparingly so in cold water. The anhydrous methiodide has m.p. 248° (Claus, Annalen, 1892, 269, 232). The anhydrous ethiodide is similar and has the same m.p. Tho dimethiodide,

$CH_2O-C_2H_2N(CH_3l)-CH(OH)-C_7H_{31}N(CH_3l)-CH-CH_2$

crystallises in yellow rhombic plates with 1.5H.O, m p. 156°. It is more readily soluble in water, but less readily so in alcohol than the monomethiodide.

Acetylquinidine,

CH,O-C,H,N-CH(O-CO-CH,)-C,H,,N-CH-CH,,

is formed on warming quantidine with acetie anhydride at 60°-80°. It is an amorphous powder readily soluble in ether (Hesse, Annalen, 1880, 205, 318).

a-Chlorohydroquinidine,

CH_O-C.H.N-CH(OH)-C,H.,N-CHCI-CH.,

is isolated as the dihydrobromide from the with 2H,O from 70% alcohol. The anhydrous base has m p. 206°, decomp. 225°, [a]21 +2763° (c=0-5 m N-HCl). The dihydrobromide has

m p. 253°, [a]p +200° in water.

The mother liquors from the crude dihydrobromide contam a'-chlorohydroquinidine, isolated as the neutral tartrate. The base crystallies from dilute alcohol in needles with 3H.O. The anhydrous base has m p. 200°, decomp. 229°, $[a]_{n}^{22} + 240.7^{\circ} (c=0.5 \text{ in N-HCl}).$ The tartrate. B, C, H, O, 11 H, O, crystallises from 309 alrohol, and when dried sinters at 152° and froths at 196° (Goodson, Lc.).

Iodohydroquinidine, better iododihydroquini-

dine (bydroiodoguinidine).

CH,O-C,H,N-CH(OH)-C,H,,N-CHI-CH, The dihydriedide, B 2HI (known in the rarly literature as " trihydriodoquinidine ") is formed when quantitine is warmed with hydriodic acid (Schnbert and Skraup), Monatsh. 1891, 12, 669; Lippmann and Fleissner, ibid, 1892, 13, 433), and has m p. 230°. The base liberated by means of ammonia crystallises from alcohol in prisms, and has m.p. 205°-206°. a-180Quinidine,

CH2O-C+H2N-CH(O)-C+H11N-CH-CH3

is formed when quinidine is warmed with strong hydrohromic or hydrochloric and for 50 hours, and the product boiled with alcoholic potash (Domański and Suszko, Bull. Acad. Polonaise, A. 1933, 119). It separates from aqueous acetone with 1H,O and melts at about 60°. The anhydrous base (from dry acctone) melts at 133° and has [a]n +111° in alcohol. It gives the fluorescence and thalleigquin tests, and on heating with snlphuric acid gives rise to β noquandine. The hydrochloride has m p. 221.

B-isoQuinidine, isoapoquinidine (B isocupreidine) methyl ether, is a stereosomeride of the preceding substance, and is formed when quintdine, a-180- or y-isoquinidine is hested with moderately concentrated sulphnric acid or by methylating isoapoquinidine (β isoenpreidine) (Pfauni, Monatah. 1911, 32, 241; Paneth, abid. 257; Konopnicki and Suszko, Bull. Acad. Polonaise, A. 1929, 340; Henry, Solomon and Gibbs, J.C.S. 1935, 966). It crystallises from moist ether or from dilute alcohol with 3H₁O. m.p. 72°. The anhydrons base has m.p. 142°. [e]n -9 7 in alrohol, or +29 in 01N. H,60, The neutral sulphate, B. H.SO, 7H.O. has [a]p -47-I' in water (when dehydrated) (Henry

et of), or -35 5° (as hydrate) Pfannl). y-EsoQuanidine is a stereoisomeride of the two preceding substances, and is formed along with a seoquinedine and with niquidine when quinidine as warmed with strong hydrobromic or hydrochloric acid for 50 bonrs, and the product boiled with alcoholic potash (Domański and Sustko, Rec. trav. chim. 1935, 54, 481). It crystalluce from aqueous acctone with $2H_2O$, mp. about 70° . The anhydrous base, which cannot be erystallised, has $[a]_D + 51 \cdot 1^\circ$ in alcohol. The acid intrate has m.p. 196° and the hydriodide, m.p. $180^\circ - 183^\circ$.

Methylapoquinidine,

$$CH_3O-C_9H_5N-CH(OH)-C_6H_{10}N>C=CH-CH_3$$

an isomeride of the preceding three substances, is fermed together with \$\beta\$-isoquinidine when quindine is heated with 60% sulphuric acid, or by inethylating apoquinidine (Henry, Solomon and Gibbs, J.C.S. 1935, 966). It separates from ether or alcohol in anhydrous crystals, m.p. 185°, $[a]_1^{15}$ +278° in 01N·H₂SO₄, or +1932° in alcohol. From aqueous acetone it crystallises with 1½H₂O, mp about 90°. The hydrochloride B·HCl, H₂O, crystallises from water or alcohol, the anhydrous salt has m.p. 267°, $[a]_D$ +174 7° in water. The geometrical isomeride of this substance, accosoquinidine, crystallises from ther with 2H₂O; it has mp. 83°, and $[a]_D$ +198 6° in dil. H₂SO₄ or +98 7° in alcohol (Henry, Solomon and Gibbs, 161d 1937, 696).

Neguidine (Domański and Suszko, Bull. Acad. Polonaise, A. 1935, 457) is claimed to be isomeric with quantiline, and to have the partial structure

$$CH_3O-C_9H_5N-CH(OH)-CH_{NH}$$

in which the quinucliding bridge has undergone rupture. It is also claimed to be an analogue of inquine (see under Quinine) the composition of which is still in dispite. Niquidine occurs along with a- and y-isoquinidine when the halogeno-hydroquinidines are dehalogenated with alcoholic potash. It crystallises from acctone, m.p. 161°, [a]_D +171° in alcohol. The acid exalate has m.p. 206°-207° and the dihydrobromide, m.p. 250°. It has a secondary nitrogen atom and forms N-acetyl and diacetyl derivatives. apoQuinidine,

$$HO-C_3H_5N-CH(OH)-C_6H_{10}N=C>CH-CH_3$$

19 one of the phenohe products formed when quinidine is demethylated with boiling 60% sulphure acid (Henry, Solomon and Gibbs, J.CS. 1935, 966). It is soluble in sodium h) drovide solution, from which it is precipitated again by earbon dioxide. It eristallises from alcohol with 1EtOH (mp. 172°) or from acctone with 1Me₂CO (mp. 178°-180°), and these can only be removed at 150°-160°. The anhydrous base has mp. 185°-190 and the second of the secon and $[a]_{0}$ +291.4° in dil H₂SO₄ or +208.6° in alcohol (figures cale for anhy drons base from measurements on the alcoholate). The hydrochloride, B.HCl, which is yellow in aqueous solution, crystallises from alcohol with 1EtOH (mp. 183°-185°) which is replaced by 2H₂O on exposure to air (mp 235°), but which is not expelled on drying at 120°. The dihydrate of this falt is also obtained by crystallisation from water: 1H2O is readily expelled at 120°, the other is more tenaciously retained. The geometrical isomeride of this substance, neoapo-

from aqueous acctone with $2H_2O$, mp. about $+120.7^{\circ}$ in alcohol (Henry, Solomon, and Gibbs, 70°. The anhydrous base, which cannot be still 1937, 595).

īsoapoQuinidine (β-isocupreidine),

$$HO - C_9H_5N - CH(O) - C_7H_{11}N - CH - CH_3,$$

is formed along with apoquinidine when quinidine or β isoquinidine is demethylated by boiling with 60% sulphurie acid (Henry and Solomon, J.C.S. 1934, 1923; Ludwiczak and Suszko, Bull. Acad. Polonaise, A, 1935, 65). It is a phenolic base soluble in alkah, and on methylation affords β-isoquinidine. It erystallises from alcohol, has m.p. 245° and $[a]_{D}-126^{\circ}$ in alcohol, or $+256^{\circ}$ in $01N\cdot H_{2}SO_{4}$. The hydrochloride has m.p. 255° and is yellow in aqueous solution. Apart from apo- and isoapoquinidine, Ludwie-zakowna, Suszko and Zwierzehowski (Rec. trav. chim. 1933, 52, 847) find also " cupreidine " among the products obtained by the demethylation of "quinidine" with boiling 60% sulphurie acid. The new phenohe base is claimed to bear the same stereo-chemical relation to cupreine as quinidine bears to quinine, i.e. that it is the phenol corresponding to quinidine, and quinidino is said to be regenerated from it on methylation. The possibility, however, of such demethylation without simultaneous isomerisation has been questioned (Henry and Solomon, J.CS. 1934, 1923; Thron and Dirscherl, J.CS. 1934, 1923; Thron and Dirscherl, Annalen, 1935, 515, 252), and it has been made probable that the Polish workers' "eupreidine" is hydroeupreidine derived from the
hydroquinino which may be present to tho extent of about 20% in commercial quinidine.

epiQuinidine, C₂₀H₂₄O₂N₂, is epimerie with quinidine about the carbinol (CH·OH) group. It occurs naturally and can be isolated from "quinoidine" (the residues from quinine manufacture) (Dirscherl and Thron, Annalen, 1935, 521, 48), but is best obtained by epimerisation of quinine or quinidine by boiling with amyl alcoholic potassium hydroxide. There is formed a mixture in dynamic equilibrium of quininc, quinidine, epiquinine, and epiquinidine. quinine being removed as neutral tartrate, and the quinidine as acid tartrate, the two epi-bases are isolated as dihydrochlorides; the epiquinine is then removed as the sparingly soluble dibenzoyltartrate. The residual epiquinidino dibenzoyltartrate is purified by recrystallisation. The base crystallises from ether in gbstening leaflets, m p. 113°, $[a]_D^{10} + 102.4$ ° (c-0 8648 in alcohol). It is very readily soluble in alcohol, but less so in ether. It is more strongly fluorescent than either quinine or quindine. The dibenzoyltartrate has m.p. 167°, and the dihydrochloride, m.p. 195°-196° (Rabe and others, Annalen, 1932, 492, 252, 258-261). Quinidine Chloride,

 $CH_3O-C_9H_5N-CHCI-C_7H_{11}N-CH=CH_2$

expelled on drying at 120°. The dihydrate of this salts is also obtained by crystallisation from water: 1H₂O is readily expelled at 120°, the other is more tenaciously retained. The geometrical isomeride of this substance, neospoquinidine, crystallises from ether or alcohol, m.p. 260°, [a]₀ +206·2° in dil. H₂SO₄ or (c-1.943 in alcohol). It dissolves readily in

alcohol, benzene or chloroform fairly sparingly toon, estimation, and detection of quinine, as in dry ether, and very sparingly in light well as of the detection of impurities therein, petroleum. The hydrochonde (Heidelberger; the reader is directed to the first three sections of and Jacobs, J. Amer. Chem. Soc. 1920, 42, 1501) crystallises from a mixture of absolute alcohol and a little dry ether in cream-coloured, woolly needles, m.p. 199°-206°, [a]25 +47 7° (c=1 017 m water). It is readily soluble in water or methyl alcohol, less readily so in chloroform and almost incoluble in acctone.

Desoxyquinidine,

CH_O-C_H_N-CH_-C,H,,N-CH=CH_

is prepared by reducing a dilute sulphine acid soltmon of quantime chloride with iron filings (Kocuigs, Ber 1895, 28, 3147; Rabe, Annalen, 1910, 373, 107; Giemsa and Halbertann, Ber 1921, 54, 1200) It separates from ether or from aqueous arctone or alcohol with 2H.O. m p 80°-82°, [a]20 for anhydrous base in alcohol +211° (Rabe) or +194° (Giemsa and Halber-Lann) It shows an intense violet fluoresrence in terteric acid as well as in alcoholic solution The hydrochloride (Heidelberger and Jacobs, J Amer. Chem. Soc 1920, 42, 1501) crystallises from a mixture of alcohol and ether with the equivalent of 1H.O, m p indef 125°-145° (dry), and [a]w +82° in weter (dry) The salt is readily soluble in water, alcohol, or chloroform, feirly readily in hot acctone and somewhat also in benzene Desovyquinidine, which elosely resembles quinene (q 1 , under Quinine), is characterised by the ready solubility of its acid tartrate Quitenidine,

CH,O-C,H,N-CH(OH)-C,H,,N-CO,H, is obtained by the oxidation of quinidine in sulphune and solution at 0° with potassium permanganate (Forst and Böhringer, Ber. 1882, 15, 1659) It crystallises from hot water

or from dilute alcohol in leaffets with 2H.O. mp 240°-246° (decomp), [a] +253° (c=5 of anhydrous aubstanco in N.H.SO.) (Goodson, llenry, and Macfie, Biochem J 1930, 24, 881) It is soluble in 200 parts of cold water, but more readily in hot waterand very sparingly in alcohol, As an acid it is soluble in alkalis, it shows the thalletoquin reaction and a blue fluoresrence in dilute sulphuric acid solution. The acid sul-

alcohol.

epiQuinidine.-See under Quinidine. Quinine, C20 H21O2N2 is the most important of the cinchona alkaloids. It was first isolated by Pelletier and Caventou (Ann. Chim 1820, 15, 318) from the "einchonino" of Gomes [1810], a crystalline mixture of alkaloids obtained from Vanquelin's "quina" (1809), the latter was an amorphous extract of cinchona bark, i.e an M/40 solution) (Buttle, Henry and and had been first prepared by Fourtroy in Trevan, Biochem. J. 1931, 28, 438). One part of 1792. Quintine occurs in all genuine emchana quintine requires for solution nearly 2,000 parts of barks, and particularly in Calisaya bark, the cold water, but dissolves in about 1,000 parts of cultivated descendant of which, C. Ledgeriana, boiling water; the presence of fixed alkalisheing made to yield as much as 8% of quinine. Ioners the solubility, but ammonia augmentest, For an account of the occurrence of quinne, supersaturated solutions being readily formed of the cultivation and commercial history of Quinno dissolves in less than its own volume of cinchona plantations, of the extraction, prepara- alcohol; in other its solubility depends greatly

this article.

The quinine of commerce usually contains cinchonidine and hydroquining as impunites These are readily removed by recrystallisation of the aulphate, acid sulphate, dahy deobromide, d camphorsulphonate, or d-beomocamphor, sulphonste, or by regeneration of the base from the periodide known as herapathite (see Hesse, Pharm. J. 1884, 15, 869; 1885, 16, 358, 818; 1886, 17, 585 , Tutin, ibid. 1909, 83, 600 : Emde, Helv. Chim Acta, 1932, 15, 574; Buttle, Henry, and Trevan, Biochem. J. 1934, 28, 433). Hydrogumine may also be removed by the mercuric arctato process of Theon and Dirsched (Annales, 1935, 515, 252).

The percentago composition of quinine (C₁₀H₁₂ON) was first correctly determined by Liebig (1838); the empirical formula

C,,H,,O,N,

assigned to it the same year by Regnault was not established until 1854, when Strecker confirmed it by analyses of quinine all yliodidis (Annales, 1854, 91, 155). For the constitution of quinine and other cinchona alkeloids, r. section 5 (p. 137) of this article Quinine hase is precipitated from aqueous sed

solutions of its selts at first es a n hite, anhydrous, amorphous powder. To avoid precipitating the frequently aparingly soluble neutral salts (particularly, for example, the sulphate.

2(B H,SO,)+2NaOH =B, H, SO, + Na, SO, +2H,O),

this operation is best performed by introducing the alkaloidal solution very slouly into e large excess of the well stirred solution of the sikeli. When emmonia is used as the precipitating agent, the amorphous base gradually turns into a crystellino efflorescent tribydrate, B3H,0, m p. 57°. 1H,O is lost on drying in the sir, so that the article of commerce is usually B 2H,0; the other two water molecules are removed over concentrated sulphuric acid or on drying in an oven The anhydrous base can also be obtained by crystallisation from hot dilute alcohol, boiling water, or ammonia solution at 100° under pressure, or by precipitating a hot solution of a quinino salt with sodium carbonate or bicarbonphate, B H2SO4.3H2O, cry stallises feom dilute ate Quinine separates from benzeno in crystals of the composition B, C, H, Anhydrous quinine base has m p. 173 5°, [a]D in alcohol -169 3° (c=2) (llesse, Annalen, 1875, 178, 206), or -167 5° (e=1 6) (Oudemans, ibid. 1876, 182, 41), or -158 2° (e=2 136) (Rabe, ibid. 1910, 373, 100); [a]_D in chloroform -116° (c=2); [a]_D in dilute hydrochloric or sulphuric acid -278 to -279° (c=16) (Oudemans), or -2815' (c=08,

on the quality of the solvent, and widely! varying figures have been published (1 in 4.5 to 1 in 114); it is more soluble in wet ether than in the dry solvent, and, as with ammonia, readily forms supersaturated solutions, which may account for some of the high solubilities On evaporation, ethereal solutions leave a residue consisting of a resinoid mass. Oninine is very soluble also in chloroform (1 in 2 parts), but requires nearly 200 parts of carbon tetrachloride. It is fairly soluble in ethyl acctate, sparingly in cold but fairly readily in hot benzene, and is almost insoluble in light petroleum (b.p. 60°) (1 in 5,000 parts). Dilute solutions of quinine in dilute sulphuric and other oxy-acids exhibit a blue fluorescence, which is weakened and may even be destroyed by the addition of halides, ferrocyanides, or thiosulphates. property enables one part of quinine to be detected in several hundred thousand parts of water. The thalleioquin reaction is not quite so delicate; a faintly acid solution of quinine is treated with chlorine or bromine water drop by drop until there is a slight excess of the halogen (bleaching powder is sometimes used), when on addition of an excess of dilute ammonia solution, a rich emerald green coloration or precipitate is produced.

Quinine undergoes a variety of chemical reactions; the products are described in greater detail below. On heating with dilute organic acids, or on melting certain salts, or on heating them in glycerol, quinotoxino (quinicine) is formed; with mineral acids quinine undergoes isomerisation to isoquinine, and also simultaneous demethylation, apoquinine and other phenolic substances being formed. Potassium permanganate oxidises it to quitenine, and chromic acid to quininic acid and to meroquinene and its oxidation products (see under Cinchonine). On catalytic hydrogenation it

gives rise to hydroquinine.

Quinine Sulphale, B. H. SO, crystallises from boiling water probably with 8H.O. Being very efflorescent, however, the water content is difficult to determine with accuracy, and is often given as 7 and 7.5H2O. The latter hydrate is official in the "British Pharmacopæia," 1932 (p. 370). The salt, the appearance of which is influenced by impurities, forms a light mass of colourless, odourless, glistening silky needles, of intensely bitter taste. They rapidy effloresee on exposure or on warming at 50° with the formation of lustreless crystals. B. H.SO4.2H.O, which constitutes the stable form, and is formed also from the anhydrous salt on exposure. The anhydrous salt, which can be obtained by drying at 100° , has $[a]_{D} - 235^{\circ}$ (c=1 in dil. $H_{\bullet}SO_{4}$) (Tutin, Pharm. J. 1909, 83, 603). The salt with $7.5H_{\bullet}O$ requires for solution nearly SO_{\bullet} parts of cold water, but dissolves in 30 parts of boiling nater. It is soluble in 65 parts of cold 90% alcohol, less soluble in absolute alcohol, and very roluble in hot 90% alcohol. It also dissolves in 30 parts of glycerol, but hardly at all in other or chloroform, although in a mixture of the latter with alcohol it is more soluble than the other solvent alone. Quinine sulphate is faintly alkaline to litmus, and on exposure to light slowly turns yellow.

Quinine Acid Sulphate (bisulphate),

B.H.SO4,7H.O,

crystallises from water in small, colourless, transparent or opaque, aci ular crystals. It is odourless, but intensely bitter. The salt is efflorescent and light-sensitive. Aqueous solutions are fluorescent, and acid to litmus, but not to Congo red. The anhydrous salt has m.p. 160° (decomp.) and [a]¹⁵ -211·7° (c=1·06, i.c. an M/40 solution in water). It is soluble in about 10 parts of cold water or 20 of alcohol, but is much more soluble at lugher temperatures. The "tetrasulphate," B·2H₂SO₄, crystallises from water with 7H₂O or from alcohol with 5H₂O. It is very soluble in water, less so in alcohol.

Quinine Hydrochloride, B·HCl,2H₂O, forms colourless glistening needles, which are efflorescent in warm or dry air. The anhydrous salt has m.p. 158°-160°, and [a]_D -155·8° in water, and about the same rotation in alcohol. It is remarkable in having a greatly diminished rotation in chloroform (-57° in 1%, and only -21° in 9% solution). The solubility of this salt in water is given variously between 2·5 and 6%; it is very soluble in alcohol, chloroform, or boiling water. The dihydrochloride, B·2HCl, forms a white crystalline powder, m.p. 180°-185° (decomp.), very soluble in water, less so in alcohol.

Quinine Hydrobromide, B·HBr, crystallises with 1H₂O according to Hesse (Chem. Zentr. 1902, ii, 953), but with 2H₂O according to the "British Pharmaceutical Codex," 1934, 887. It is soluble in 55 parts of cold water, but is readily soluble in alcohol, chloroform, or boiling water. The dihydrobromide, B·2HBr,3H₂O, is very soluble in water or alcohol.

A very large number of other salts, derivatives and preparations of quinine, are known; only a small selection, however, can be given here of those that are available for use in

medicine.

Quinine and Urea Hydrochloride,

B-HCI-CO(NH.).-HCI,5H.O.

is prepared by dissolving a molecular proportion of urea (60 parts) in a solution of quinine hydrochloride (400 parts) in 300 parts hydrochloric acid (sp.gr. 1.05). The salt having the above composition crystallises out. It has m.p. 72° and is soluble in one part of water and two parts of alcohol. It is acid to litmus in aqueous solution, and gives the fluorescence and thalleioquin reactions characteristic of quinine. A similar preparation is quinine and urethane hydrochloride; usually prepared as a solution for injection ("British Pharmaccutical Codex," 1934, pp. 889, 1287).

Quinine Acetylealicylate, B.C₂H₈O₄, prepared by mixing alcoholic solutions of the components, is a white crystalline powder, m.p. about 160°, soluble in 330 parts of water, or 50 of

alcohol.

Quinine Citrate, B₃·C₆H₈O₂.7·5H₂O, or B₂·C₆H₈O₂.7H₂O, white needles, soluble in about 1,000 parts of water.

Ouinine Disalicylosalicylate or bisalicylo- | salicylate (" quinisal," " quinisan ").

It is described as a white microcrystalline powder baving only a slightly bitter taste, insoluble in water and very soluble in alcohol. Quinine Formate (" quinoform "), B CH,O.

consists of crystals, m p. about 130°, fairly soluble in water.

Quinine

Glycerophosphate ("Lincurine"), B. C. H.O. H. PO. 4H. O. from quinine hydro chloride and calcium glyrerophosphate, occurs in crystals, m p. 154°, soluble in 200 parts of water. It is bitter.

Quintne Hypophosphile, B H2PO2,2H2O, 25 a crystalling or amorphous powder fairly readdy soluble in water.

Quinine Iodobiemuthale, (BI(1), B 2H1, a red amorphous powder prepared by mixing solutions of quinine by drocbloride, bismuth chloride, potassium iodide, and hydrochlone acid (Fran-cols and Seguin, J Pharm Chim, 1925, 2, 59).

Quiname Phosphate, B, 2H, PO, 6H, O, or B, H, PO, 8H, O, occurs in needles which resemble quinine sulphate both in appearance and solubility

Quinine Salicylate, B C, H,O3,H2O, from quinine sulphate and sodium salicylate in aqueous solution. It occurs in white silky needles, very sparingly soluble in water

Quinine Tannate is prepared by mixing solutions of quinine sulphate and tannic seid in the presence of ammonia It is a pale yellow, amorphous powder, having an astringent, harely bitter taste. It is slightly soluble and only slowly absorbed.

Quinine Valerale (valerianate),

consists of colourless, lustrous, pearly crystals, or of a white, micro-crystalline powder, m.p. about 90°, slightly soluble in water, and having a bitter taste and a faint odour of valerie seid. Quinine Carbonate (diquinine carbonate, "ari-

stochin "). CH.O-C.H.N-CHO-C.H.,N-CH-CH.

is a tasteless, amorphous, insoluble powder, m.p. 189° made by treating quinine with phosgene in benzene solution.

Quinine Ethylearbonate ("British Pharmacoperta," 1932).

CH,O-C,H,N-CH(O-CO-OC,H,)-C,H,,N-CH-CH,. occurs in white, almost tasteless, needles, m.p. sbout 90°, slightly soluble in water. It is made by treating quinine with chloroformie ester (Cl CO₂C₂H₂) in benzene solution.

The following are the principal derivatives, transformation, and degradation products of quinine, which at present are only of academic interest :

Acetylouinine.

CH, O-C, H, N-CH(O-CO-CH,)-C, H, IN-CH = CH, was first prepared by Hesse (Annales, 1880, 205. 317; cf. Beckett and Wright, J.CS 1876, 29. 657) by warming quinine with acetic anhydride Other methods bave been patented by Vereinigte Chimafahriken, Zimmer & Co. Gmb.if (GP. 128116/1902, 178172, 178173/1906). 1 crystallises from ether in prisms, m p. 103°, or from light petroleum, m p. 116°-117°, bas [a]n -54 3° in alcohol, or -114 8° in dilute acid, and

is readdy soluble in alcohol, ether or chloroform It is almost tasteless and can be bydroly sed back to quining,

Benzoylyuinine,

CH_O-C_H_N-CH(O-CO-C_H_)-C_H,1N-CH-CH, was first made by Schutzenberger (Annales, 1858, 108, 352; cf. Wunseb, Ann. Chim, 1896, [vii], 7, 127) by the action of excess of benzoyl chloride on quipine on the water bath. Other methods have been patented by Vereinigte Chmufabriken, Zimmer & Co., Gn (GP 128116/1900, 178172, 178173/1905) crystallises from ether, mp 139°, [a]0 +121 6' in alcohol, and forms a hydrochloride,

B HCI, HO,

which when dry bas m,p 226°. Quinine Methiodide,

CH2O C2H4N CH(OH) C7H11N(CH3I) CH CH3 is obtained when equimolecular proportions of the components react in etheresl or alcoholic solution in the cold (Claus and Mailmann, Ber. 1881, 14, 76; Hesse, stid 1895, 28, 1298) It ervatalises from water in colourless needles. with 1 or 2H.O, mp 233°-236° (decomp), is very sparingly soluble in ether or chloroform,

sparingly in cold water, but readily so in alcohol. The dimethiodide, CH,O C,H,N(CH,I) CH(OH) C,H,,N(CH,I) CH CH,,

is formed from it (also from cupreine and from quintie) by boiling with an excess of methyl jodide in alcoholic solution. It crystallises from water in yellow plates with 3H,O, mp about 160° (decomp.). Quinine Ethiodide,

CH,O C,H,N CH(OH) C,H,,N (C,H,I) CH CH,

erystellass from water and IM, O. The ar hydrous substance has m p. 210°-211° and [s], -105° m slcobol. It is readily soluble in alcobol, but sparingly so in rold water.

Isomeric Quinine Ethiodide,

CH'O C'H'N(C'H'I) CH(OH) C'H'IN CH CH' (Skraup and von Norwall, Monatsh. 1891, 15, 47)

is obtained as the hydricdide (m p. 231°, dry) hy heating quinine hydrodide with ethyl iodide in alcoholic solution at 100° (scaled tube). It crystallises from 50% alcohol in needles with 3H.O, m p. 93°, and is very soluble in alcohol, hut spanngly so m water.

Quinine Diethiodide,

CHIO C.H.N(C.H.I) CH(OH) C.H.IN(C.H.I) CH CH, prepared from either of the two preceding substances by bothing with ethyl jodicie in alcoholic solution, ery stallines with 1H,O, m p 157°-167.

a-Chlorohydroquinine, better, a-eblorodihydroquinine (component of "hydrochloroquinine"),

CH2O·C2HEN·CH(OH)·C7H11N·CHCI·CH2,

is prepared either by leaving quinine hydrochloride to stand in hydrochloric acid (saturated at -17°) for several weeks (Comstock and Koenigs, Ber. 1887, 20, 2517), or by heating with the ordinary concentrated acid at 85° in a scaled tube (Hesse, Annalen, 1893, 276, 125). a-Chlorohydroquinine is isolated from the products and purified first as acid nitrate, and then by crystallisation and recrystallisation of the base from methyl alcohol (Goodson, J.C.S. 1935, 1095), when it separates in anhydrous rhombs, m.p. 210°-215° (decomp.), [a]_D²⁰ -251° (c=0.5 in N-HCl). The acid nitrate B.2HNO₃, has m.p. 212°, and the tartrate, B₂·C₄H₆O₆·7H₂O₇·198° (dry).

a'-Uhlorehydroquinine is epimeric with the

preceding substance at the earbon atom which carries the chlorine atom. It is formed along with it, the mixture constituting "hydrochleroquinine" (Goedson, I.c.). It is found in the mother-liquors from the methyl alcoholic erystallisation of the bases, whence it is isolated as acid nitrate, the base recovered, and recrystallised from benzene. It crystallises with solvent of crystallisation, but on exposure to air, the benzene is replaced by water. The anhydrous base has m.p. 194°, decomp. 225°, The annyarous was non-zero in N-HCl). The and $[a]_D - 168\cdot 1^\circ$ (c=0.5 in N-HCl). The acid nitrate, B·2HNO₃, has m.p. 223°, and the lattrate, B₂·C₄H₆O₆·2H₂O, 228° (dry).

a.Bromohydroquinine, a-bromodihydroquinine, hydrobromoquinino II (component of "hydrobromoquinine "),

CH3O·C,H&N·CH(OH)·C,H11N·CHBr·CH3,

is prepared by eausing quinine dihydrohromide to react with hydrohromic acid (saturated at -17°) at room temperature for 3 days (Comstock and Koenigs, Ber. 1887, 20, 2518), or by heating quinine hydrochloride with 38% hydrobromic acid in acetic acid at 80° for 21 hours (Podlewski and Suszko, Rec. trav. chim. 1936, 55, 392). a-Bromehydroquinine hase is isolated from the products by crystallisation from ether, and purified by recrystallisation from acctone, when it reparates in the form of prisms, m.p.166 °-167° (decomp.), $[a]_D -200^\circ$ in alcohol-chloroform (1:2). With alcoholic potash it gives rise to niquine (see below)

a'-Bromehydroquinine (hydrohromoquinine I), epimeric with the preceding substance at the carbon atom which carries the bromine atom (Podlewski and Suszko, I.c.), is formed along with it, and is isolated from the ethereal motherliquors of mixed bromo-hases by recrystallisation from benzene. It separates with one molecule of solvent of crystallisation, has m.p. 160°-162° (decomp.), and $[a]_p - 50^\circ$ in alcohol-chloroform. With alcoholic potash it gives rise to β-isoquinine (see below). Its quarternary metho-p-toluenesulphonate crystallises from water, m.p. 193°-

1910

a-Iodohydroquinine. a-iododilydroquinine, hydroiodequinine II (component of "hydroiodoquinine ''),

CH3O-C3H5N-CH(OH)-C7H11N-CHI-CH3,

is formed when quinine is heated with hydriodie acid (sp.gr. 1.7) on the water hatb for 1 or 2 hours (Lippmann and Fleissner, Monatsh. 1891, 12, 327, and later papers; Skraup, ibid. 431, and later papers). The epimerides formed isolated as dihydriodides; the bases are then liberated, and separated from one another by erystallisation from benzene and from ether (Rosenmund and Kittler, Arch. Pharm. 1924, 262, 20; see also Reyman and Suszko, Bull. Acad. Polonaise, A, 1935, 360). a-Iodohydroquinine on heating begins to decompose above 100°, and the decomposition reaches a chimax at about 150°. The specific rotation is—128° (Rosenmund) or —217° (Reyman and Suszko). On boiling in benzene selution, or on treatment with alcoholic potash, niquine (see below) is

a'-Iodohydroquinine, hydroiodoquinine I, is the epimeride of the foregoing compound. According to Rosenmund and Kittler (l.c.) it has the same melting-point, but $[a]_D -74.3^\circ$. According to Reyman and Suszko (l.c.), however, it crystallises with one molecule of benzene and melts at 93°, re-solidifies and then decomposes at 130°-140°; it cannot be freed from benzene without decomposition; the crystals containing benzene bave $[a]_D -18^\circ$ in alcohol. The methiodide has m.p. 112°-114°. On boiling with benzene, or on treatment with alcoholic silver nitrate or potash, a'-iodohydroquinine gives rise to β -isoquinine (see below).

The hydroxy-compound corresponding to the above halogen-derivatives is said to be formed when quinine is treated with cold concentrated sulphuric acid, and the resulting sulphonic acid saponified by heiling with dilute sulphuric acid (Vereinigte Chininfabriken, Zimmer & Co.,

G.m.b.H., G.P. 152174/1904).

By treating quinine with the halogen hydracids and with sulphuric acid under more drastic conditions, there is obtained a series of halogenohydroapequinines and a hydroxyhydroapequinine respectively, differing from the proceding substances only in having a hydroxyl in place of the methoxyl group in the 6-position of the quinoline nucleus. These plienels, however, have not been so fully studied.

a-isoQuinine, isoapoquinine methyl ether (component of "pscudoquinine"),

 $CH_2O-C_9H_8N-CH(OH)-C_6H_{10}N>C=CH-CH_3$,

is formed in small quantity when quinine is heated with sulphurie acid (sp.gr. 1.61 at 100°) (Bötteher and Horovitz, Monatsh. 1912, 33, 571), but is probably better obtained by the methylation of iscapequinine. It is isolated from the other products of the reaction as the tartrate, and is purified by recrystallisation of tho base from benzene. It has m.p. 196.5°, [a]D -245° (c=1 in alcohol).

β-isoQuinine, isoquinine, apoquinine methyl ether (component of "pseudoquinine") is the geometrical isomeride of the preceding substance. It is formed together with a-isoquinine, niquine, and other substances when quinine is heated with sulphurie acid (sp.gr. 1.61) at 100° (Bötteher and Horovitz, ibid. 576), when the halogenohydroquinines are dehalogenated (see, example, Suszko, Bull. Acad. Polonaise, A, 1925,

with Gibbs, ibid. 1935, 968). In the first two cases it is formed as the principal component of the products, which may contain also unchanged bromide (Henry and Solomon, J.C.S. 1934, quinine, a isoquinine, niquine, and hydroxyhydroquinme. It is isolated together with the quinine and a-isoquinine in the ether soluble fraction ; the quinine and a isoquimme may be removed as the snlphate and tartrate respectively, the β-isoquinine being then isolated as the oxalate, and purified by recrystallisation of the base preferably from arctone, but also from dilute alcohol, ether or benzene It has m p. 189° (Böttcher), [a]p -201° in alcohol (Henry) or -102° (-195°) (Bottcher, Sasako), -295 9° in dilute sulphure acid. The hydrochloride has mp. 249°-251°, [e]_D -174° in water, and the sulphate, mp 220°.

Niquine, of disputed composition, the formulæ in question being C₁₉H₂₄O₂N₂, C₂₆H₂₂O₂N₂, and C₂₀H₃₄O₂N₃, is formed when todo, or bromoh) droquinme is dehalogenated. This may be effected either by treatment with alcoholic potash or alcoholic silver mitrate, or by boiling the halogeno bases in benzene solution. When using the crude mixture of a- and a' halogeno formed by the treatment of cumme with halocen hydracids, the dehalogenation gives rise to a mix ture of niquine and B-1400 unine. It is also said to be formed in minute amount together with β seequanne, when quante is treated with sal phuric acid (Giemsa and Gesterlin, Ber 1931, 64, [B], 60) Niquine crystallises from boiling water, or dilute alcohol, with 2H,O, m p. about 100° It is almost insoluble in moist ether. The water is lost in a desiccator and the anhydrous base then melts at 146°, and becomes soluble in dry ether. The anhydrous base has (a)_D -1°2° in alcohol. It is generally agreed that inquine contains the hydroxyl group of quinine intact , but that the quinuclidine nitrogen atom is changed from tertiary to secondary, because miquine forms a diacetyl-derivative, and a Nnitrosamine. It reacts additively with hydrodic acid, and is largely regenerated from the resulting compound by treatment with alcoholic potanh, a small amount of "ssoniquine" being also formed (Skraup, Monatsh, 1893, 14, 440; for the remaining literature of miquine, see Lippmann and Fleissner, ibid. 555. Rosenmund and Kittler, Arch. Pharm. 1921, 262, 23; Suszko, Bull. Acad. Polonaise, A. 1925, 129. Reyman and Suszko, ibid. 1935, 360; Podlewski and Suszko, Rec. trav. chim. 1936, 55, 392; foc constitution by analogy with cauchonhydrine, which will be found under Cinchonine, see also references given there).

apoQuinine, \$ 150cupreine,

HO-C,H,N-CH(OH)-C,H,11N> C+CH-CH,, is the principal crystalline component of the amorphous mixture obtained when quinne is demethylated or cupreme isomerised with hot dilute mineral acids or with alumining chloride. The amorphons matture was first obtained by Hesse (Annalen, 1850, 205, 223) who, takings to be a single substance, called it "a programme for the proper of the proper of

129), and when a poquinine is methylated (see, for Suszko, Rec. trav. chim. 1933, 52, 841), and can example, Henry and Solomon, J.C.S. 1934, 1928; be freed from its congeners by repeated reprecipitation, followed by fractional crystallisa-tion of the acid sulphate or of the dihydro-1927). opoQuinine, which is persistently amorphous until fairly pure, crystallises from ether. alcohol or arctone, in each case with solvent of erystallisation, which is removed with difficulty, and not without loss of the crystalline state. It froths at 184°, and the froth becomes a clear liquid at about 205°; the anhydrous base has [a]20 -214 8° (c=0-775 in alcohol), or -281° (c=0.784 in 0.1 N.H. 201). apo-Quinine is a phenolic substance, soluble in alkalis and is precipitated therefrom by passage of carbon dioxide. In common with other phenolic bases of the cinchona series, it forms neutral salts, which are intensely yellow m aqueous solution, the colour being discharged by the addition of acids. It is non fluorescent, and does not respond to the thallelogum test. A series of allyl ethers has been prepared in Europe, America, and in Japan, which promise to be of similar interest to the corresponding ethers of hydrocupreme (see Buttle, Henry, Solomon, Trevan and Gibbs, Biochem. J. 1933 The methyl ether is \$ isoquinine, 32, 47). apoQuinine hydrochloride crystallises from water or alcohol, m.p. 272 5°, [a]p -163 8° in water. The acid sulphate, when anhydrous, has [a]n -223 in water; the sesquiordist melts at 225, and the dihydrodromide, which crystallness from alcohol, mp. 235, [a], -180-9 in water, tends to lose HBr by dis-

sociation isoapoQuinine, a-isocupreine, is the metrical isomeride of apoquinine, which it greatly resembles, and on methylation yields a sequence. It is formed together with apo-quinine and hydroxyhydroapoquinine when quinine is demethylated with sulphuric and (Henry, Solomon and Gibbs, J.C.S. 1935, 967) It occurs in the mother-liquors from the and sulphate or dihydrohromide fractionations of impura apoquinine. The latter salt, hewever, although the more soluble of the two in alrohol, is the less soluble in water. The base crystallises from arctone containing a little alcohol with the equivalent of 3H,O, or preferably from meth)! alcohol. It froths at 190° and maily derom-poses at 275°. The substance, like a reoquining. is remarkable for its high optical rotation, the anhydrous base having [a]15 -261-70 (c= 0-775 m alcohol), or -364 m dilute acid. The hydrochloride has m.p. 271° and [a]p -1945° in water; the dihydrobromide has m.p. 136°-140°, being thereby readily distinguished from opoquinine dihydrobromide, which melts 2t

97.50

Ournotoxine (quinteine). CH.O.C.H.N.CO.CH, CH, CH, N.CH - CH,

is said to occur in canchonn back, but is best

Ber. 1900, 33, 3228). It has also been obtained by heating quinine with glycerol at 210° (Hesse, Annalen, 1873, 160, 277). Quinotoxino is a yellowish amorphous mass, which becomes an oil at about 50° to 60°. It has [a]n +44·1° (c=2 in chloroform) (Hesse), or +38.7° (Howard and Chick, J.S.C.I. 1909, 28, 55). It is almost insoluble in water, but readily soluble in alcohol, ether or chloroform. It is a strong base which absorbs carbon dioxide from the air, and dissolves, when freshly precipitated, in ammonium salt solutions, expelling ammonia. It dissolves in mineral acids with a yellow colour, but does not fluoresce. With chlorine water and ammonia it produces the characteristic green coloration (thalleioquin). Quinotoxine hydrochloride (B.HC1) is recommended for the isolation of the toxine; it has mp. 180°-182°, [a]_D +15° (average) in water (Heidelberger and Jacobs, J. Amer. Chem. Soc. 1919, 41, 832). The neutral sulphate, B2·H2SO4,3H2O, erystallises from alcohol in prisms, and dissolves readily in water. The ozalate, B₂·C₂H₂O₄,9H₂O, often used for the isolation of the toxine, erystallises from chloroform or alcobol, and is very sparingly soluble in water; the anhydrous salt has m.p. 166°-167°, [a]_D²⁰ +24° (c-2 ln 1 vol. alcohol and 2 vols. chloroform) (Dirscherl and Thron, Annalen, 1935, 521, 64). Other erystalline salts are the bydriodide, thioeyanate, tartrate, and platinichloride. Quinotoxine is a ketonic secondary-tertiary base. Its p-bromophenylhydrazone melts at 141°, and its Nomethyl-derivative, identical with the "methylquinine" of Claus and Mallmann (Ber. 1881, 14, 79) and the "methylquinidine" of Claus (Annalen, 1892, 269, 234), can be obtained either by direct methylation or by boiling quinine or quinidine methodido with aqueous alkali (phenylhydrazone, m.p. 135°-136°). Quinotoxine forms an isonitroso-derivative with amyl nitrite, and a N-nitroso-derivativo (nitrosamine) with nitrons acid.

epiQuinine, C20H24O2N2, is epimeric with quinine about the earbinol (CHOH) group. It occurs naturally, and can be isolated from "quinoidine" (Dirscherl and Thron, Annalen, 1935, 521, 48), but was first obtained by epimerisation of quinine or quinidine by boiling with amyl alcoholic petassium hydroxido when there is formed a mixture in dynamic equilibrium of quinine, quinidine, epiquinine and epiquinidine. The quinine and quinidine being removed as neutral and acid tartrate respectively, the two cpi-bases are isolated together as dihydrochlorides; the epiquinine is then separated as the sparingly soluble dibenzoyltartrate, which cannot be recrystallised, and can only be purified by recovering the base and converting back to dibenzoyltartrate. The pure base is a colourless, viscous oil, [a]_D²² +43.3° (cm0 9494 in alcohol). It is readily soluble in most organic solvents, and is more strongly fluorescent in dilute sulphurie acid than the parent alkaloids. Its dihydrochloride erystalhecs from acctone, m.p. 196°, [a]p +33.3° in alcohol, and its dibenzoyltartrate, also from acctone, has m.p. 159° (Rabe et al., Annalen, 1932, 492, 252.)

952, 492, 252.) Vol. 111.—12 Quininone.

 $CH_2O-C_9H_5N-CO-C_7H_{11}N-CH=CH_2$ is obtained in low yield by the gentle oxidation of quinine or quinidine with chromic acid (Rabe and Kuliga, Annalen, 1909, 364, 346, 349). It is best prepared indirectly from quinotoxine by N-bromination with sodium hypobromite, followed by treatment of the N-bromotoxine (m.p. 123°), with sodium ethoxide (Rabe, Z. angew. Chem. 1913, i, 543; Rabe and Kindler Ber. 1918, 51, 466). It separates in almost colourless crystals from ether, melts at 101° (108°), and shows mutarotation, the final value at 20° being +75.5° (c=2 in alcohol) (Rabe, Kuliga and Marseball, Annalen, 1910, 373, 116, 117). It is readily soluble in most organic solvents, but not in water. It dissolves slightly in aqueous alkalis with a yellow colour (enolisa-The hydrochloride, B.HCl, forms a very hygroscopie pale yellow, erystalline powder, m.p. 210°-212°, [a]_D (final value) +58.5° in alcohol. Quininone forms a glassy, oxime and on reduction in a suitable manner gives rise to quinine and quinidine (Rabe and Kindler, l.c.). With amyl nitrite and sodium ethoxide it is split into quininie acid and oximinovinylquinuelidine.

Quinine Chloride,

 $CH_3O-C_9H_5N-CHCI-C_7H_{11}N-CH=CH_2$

is prepared from anhydrous quinine hydroehloride, chloroform and phosphorus pentachloride (Comstock and Koenigs, Ber. 1884, 17, 1988; Rabe, Annalen, 1910, 373, 103; Gienisa and Halberkann, Ber. 1921, 54, [B], 1192). It erystallises from benzene, or benzene and ether, in magnificent colourless crystals, m.p. 151°-152°, [a]_D +60·3° to +62·6° in alcohol. Quinine bromide (C. F. Böbringer & Soeline, G.m.b.H., Rabe, Cohausz and Scheel, G.P. 592541/1932) has m.p. 154°.

Desoxyquinine,

 $CH_3O-C_9H_5N-CH_2-C_7H_{11}N-CH=CH_2$

is prepared by reducing a dilute sulphuric acid solution of quinine chloride with iron filings at room temperature (Koenigs, Ber. 1896, 29, 372; Rabe, Annalen, 1910, 373, 107; Giensa and Halberkann, Ber. 1921, 54, 1197). It erystallises from moist ether or from aqueous alcohol or acetone in colourless, efflorescent needles, B·2H₂O, m.p. about 50°. The anhydrous base, which is not readily obtained crystalline, has m.p. 94° and [a]_p about +99° in alcohol. It is fluorescent in dilute sulpburic acid and shows the thalleioquin reaction.

Quinene,

CH₃O-C₉H₅N-CH=C=C_eH₁₀N-CH=CH₂, is prepared by boiling quinine or quinidine cbloride with alcoholic potash (Comstock and Koenigs, Ber. 1884, 17, 1989; 1885, 18, 1223; Giemsa and Halberkann, *ibid.* 1921, 54, 1192). It erystallises from ether, ligroin, or aqueons acctone with 2H₂O, m.p. rather indefinite between 67° and 90°. The anhydrous base has [a]_D +52·5° in alcohol. The dihydrochloride, B·2HCl, H₂O (Heidelberger and dacobs, J. Amer. Chem. Soc. 1920, 42, 1501), crystallises from a mixture of alcohol and ether in lemon-

yellow needles; the anhydrous salt has mp. | yellow colour) and in alkalis. Alcoholic solutions

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1802-185°, and [a]p +18 4° in water. apoQuinene, 4 (6 hydroxy-3 4-diethylphenyl)-6 hydroxy-quinoline,

HO-C.H.N-C.H.(OH)(C.H.)

(Kenner and Statham, JCS, 1935, 299), is obtained by heating quinene with hydrobromic acid (sp gr. 1-49) at 180° (Comstock and Koenigs, Ber. 1885, 18, 1226; Koenigs, J pr. Chem 1900, 61, 41) apoQuinene, which should not be confused with apoquinine, crystallises from aqueous alcohol, m p 246°-248°.

Quininal, CH3O-C,H3N-CH(OH)-C,H11N-CH=O, and its acyl (acetyl and benzoyl) derivatives,

CH,O-C,H,N-CH(O-CO-R)-C,H,N-CH-O, are obtained from quinine and from its acyl derivatives by the action of ozone in rold chloroform solution. The resulting ozonides furnish on hydrolysis formaldehyde, hydrogen perovide and the aldehydic substances in question. The latter are amorphous Quininal has [a]n - 30" in chloroform, the acetyl deriva tive has [a]D -63 5°, and the benzoyl derivative +79 4° (Seekles, Rec trav. chim 1923, 42, 69) Quitenine,

CH_O-C_H_N-CH(OH)-C,H,,N-CO_H is obtained by oxidising a cooled solution of

quimine in dilute sulphuric acid with potassium permanganate. It was first prepared by Kerner in 1869, and has been studied by Skrsup (Ber. 1879, 12, 1104, Annalen, 1879, 199, 348; Monatsh 1889, 10, 40) It crystalises from dilute alcohol in prisms with 4H,O, m p. 286° (decomp), [u]_D -142 7° (c≈01 m alcohol of sp gr. 0 958) The anhydrous substance has [a]p -298° (e=5 m N.H.SO.) (Goodson. Henry and Macfic, Biochem J. 1930, 24, 881) It is insoluble in ether or water, and sparingly soluble in alcohol, aqueous alcohol, or ammonia, it is readily soluble in acids or fixed alkahs, but from the latter is precipitated again by absorption of earbon dioxide from the air Quitenine is fluorescent both in acid and in aqueous-alcoholic solution and responds to the thallesoquin test On oxidation with chromic acid it gives rise to the same oxidation products (but no meroquin enc) as result in the like oxidation of quimine On demethylation with hydrodic and it gives rise to the phenolic aubstance "quitenol" ("cupretenine") (von Bucher, Monatah 1893, 14, 601), but suffers no other change. Quitenine gives an acetyl derivative, and forms salts both with mineral acids and with metals. The ethyl ester has m p 200°.

Quininic And, 6 methoxy quinohne-4 carboxylic acid, CH,O-C,H,N-CO,H, is the most readily replated of the exidation products of quinine, quinidine, their hydro-derivatives, and many of their derivatives and degradation products. It has been synthesised by Rabe and co-workers (see section 5b, p. 144a). The following properties are mainly from Skraup (Monatah 1881, 2, 589), it crystallises from ddute hydro chloric acid in yellowish prisins, in p. 280°, soluble in about 100 parts of boiling alrohol, almost insoluble in water, ether or benzene. It is readily soluble in mineral acids (with a acid

of the acid show a blue or violet fluorescence, and at responds, but not in typical fashion, to the thallesogum test. On demethy lation it gives rise to xanthoquinme acid, and on oxidation with alkaline permanganate to 234-pyridinetricarbox he acid. It forms a hydrochloride, and

salts with barrom, calcium, copper, and silver Its methyl ester has m p. 85°, and ethyl ester 69°. 6 Methorylepidine, 6 methoxy-1 methylquino-hne, CH₂O—C,H₂N—CH₂, has been obtained by the hydrolysis of quineno (Koenigs, Ber 1890, 23, 2673; Kaufmann, told. 1913, 46, 1829) and has been synthesised from anisidine by Pictet and Misner (ibid, 1912, 45, 1802). and by Rabe and co workers (thid 1931, 64, 2492). It crystallises from boiling water or aqueous alcohol with 1H2O, has m p 50°-52° h p 213° and forms a picrate, m p. 223°, and a sulphate insoluble in alcohol. It shows a blue fluorescence in water. On demethylation with boding hydrobromic acid it gives rise to p hydroxylepidine.

ebrOulnine - See under Qumme Quinoidine is the name originally applied by Serturoer (1828), the discoverer of morphine, to the amorphous material extractable from the mother liquors left after removing quinine from the total alkaloids of cipchona bark The name quinoidine is still applied to residues from quining manufacture, but as this now differs greatly from the method used by Serturner, and as the other crystalline associates of quipine are now also recovered, the material which goes by this name to-day differs greatly from Serturner's product. It contains a small proportion of quinotoxine, but consists largely, as has been shown recently, of epiquinine and epiquinidite. which may be accompanied by their dibidroderivatives (Dirscherl and Thron, Annales, 1935, 521, 48; this paper also summarises the history of quinordine). Two views have been held as to the origin of these amorphous bases; (a) that they actually occur in the bark, (b) that they originate by decomposition of the crystal line alkalords during manufacture, Dirscherl and Thron are of opinion that the epi bases at

least pre exist in the bark.
Quinotoxine — See under Quinine, (p. 176d)
"Totaquina."—See p. 132

A. M. and W. R. CINCHONAMINE r CINCHONA ALKA-LOIDS) (this vol., p. 152a).

CINCHONICINE (Cinchologine) r. Civ-CHO'A ALKALOIDS (this vol. p.157d). CINCHONIOINE v. CINCHONA ALKA-

LOTO3) (this vol., p. 1526). CINCHONIGINE (B 1soCinchonine) v CINCHONA ALKALOIDS (this vol., p. 156c)

CINCHONILINE (d. 150Cinchonine) v. Civ. CHOYA ALKALOIDS (this vol., p. 1566).

CINCHONINAL r. CINCHONA ALKALOIDS

(this vol., p. 158d)
CINCHONINE C. CINCHONA ALKALOIDS (this vol. p. 1545)
CINCHONINONE r. CINCHONA ALEA-

CINCHOPHEN PHENYLCINCHO-NINIC ACID 2-pheny lquinoline-4 carboxylic

1.01D3 (this vol., p. 159a).

CINCHOTICINE v. CINCHONA ALKALOIDS

(this vol., p. 163c). CINCHOTINE,

DIHYDROCINCHO-NINE (Hydrocinchonine) v. Cinchona Alka-LOIDS (this vol., p.).
CINCHOTINE v. CINCHONA ALKALOIDS

(this vol., p. 160a)

CINCHOTINETOXINE

Alkaloids (this vol., p. 162d). CINCHOTOXINE v. CINCHONA ALKALOIDS (this vol., p. 163c). 1:4-CINEOLE.

1:4-Cincole, C₁₀H₁₈O, b.p. 172°, d¹⁶ 0.9010, $n_{\rm D}^{16}$ 1.4479, is present in small amount in the essential oil from the fruits of Piper Cubeba, grown in Mysore (Rao, Shintre and Simonsen, J.C.S.I. 1928, 47, 92). It has been shown by Austerweil (Bull. Soc. chim. 1928, [iv], 45, 862) to be present also in the low boiling fractions of the oil obtained in the manufacture of a-terpineol. 1:4-Cincole was first prepared by Wallach (Annalen, 1907, 356, 204; 1912, 392, 62) by the dehydration of trans-terpinene-terpin with oxalic acid. It can be obtained in a similar manner from cis-terpinene-terpin and from terpin hydrate (Dupont, Levy and Marot, Bull. Soc. chim. 1933, [iv], 53, 393). With hydrogen chloride 1:4-cincole yields terpinene dihydrochloride.

1:8-CINEOLE.

The oxide, 1:8-cincole, C₁₀H₁₉O, known also as eucalyptol and cajuputol, m.p. 3°, b.p. 174-4°/ 760 mm., d^{15} 0.930, $n_{\rm D}^{15}$ 1.45839, is one of the most widely distributed of essential oil constituents. It occurs, inter alia, in many encalyptus oils, in Oleum cina (oil of wormseed) from Artemesia maritima, and in oil of cajuput from Melaleuca minor. It finds considerable use in pharmacy.

Cincole (1) can be obtained by the dehydration of terpin (11), whilst on treatment with hydrogen chloride it yields dipentene dihydrochloride (111), reactions which establish its structure. Cincole is extremely stable and distils unchanged over rodium. On oxidation with potassium permananate it gives cincolic acid (1V), m.p. 2042-

CINCHOTENINE v. CINCHONA ALKA-| m.p. 104.5°-106°. This latter acid is converted into a-cinenic acid (VI), m.p. 83°-84°, when heated at 150° (Rupe, Ber. 1908, 41, 3955; Helv. Chim. Acta, 1933, 16, 505; 1934, 17, 98).

Various methods have been suggested for the quantitative estimation of eincole. The two most satisfactory are the preparation of (i) its erystalline additive compound, C10H18O, H3PO4, which is formed when it is mixed with syrupy phosphoric acid or (ii) its compound with ocresol, C₁₀H₁₆O,C₂H₈O, m.p. 55·8° (Coeking, Pharm. J. 1920, 105, 81).

CINNABAR. Native mercury sulphide. (HgS), and the only ore of this metal. Its erystals are rhombohedral, with perfect cleavages parallel to the faces of the hexagonal prism; they are of interest crystallographically in presenting the same type of trapezohedral symmetry as quartz, and, like this, they rotate the plane of plane-polarised light. They are bright-red in colour, often transparent, and have a brilliant adamantine to sub-metallic lustre; the scarlet streak is characteristic. Sp.gr. 8-1; hardness, 2-21. The refractive indices are high: for the ordinary ray 2-81429, and for the extraordinary ray 3:14344 for red light $(\lambda 672 \ m\mu)$; and $n_E 3.27188$ for sodium-light at 21°C. (11. Rose, Centr. Min. 1912, 527). The manganate it gives cincolic acid (IV), m.p. 2042— rotatory power is also very high (over 25 times 206° (Wallach, Annalen, 1888, 246, 268; 1890, 258, 319; 1892, 271, 20), which when heated with water at 150° gives cinogenic acid (V), 186°C. (H. Rose, N. Jahrb. Min, 1900, 29, 94). The massive ore, which may be compact or with zine-copper couple (Thirle, Ber. 1899, 32, earthy in texture, is of the same colour; but 1299) yields the dimeric product hydrocinas sometimes it is liver-brown or black, owing to main, Ph-DC HC HC HOHOH, but it may be the admixture of clay or organic matter, as made to cumamyl alcohol with iron and series on the varieties called represented by Applic and (1241), Chimilt and Bohme, Ber. 1924, and the company of the compan and no mineral containing mercury has yet this reaction is very sensitive to added impurity been found in the British Isles. It is mined at and by suitably adjusting the ronditions either Almaden in Spain; Monto Amiata in Tuscany; Idria in Carmola; New Almaden in California; in the provinces of Kwei-chou and Hunan, in Soc. 1925, 47, 3061); electrolytic reduction also central China; and in Mexico and Peru.

Metac naubarate is the tetrahedral cubic modifiention of HgS, of rare occurrence as small black crystals Cinnabar and two other crystalline modifications of mercuric sulphule have been obtained artificially by E. T. Allen and J. L. Crenshaw (1912) Cinnabar is the stable form at all temperatures up to its sublimation point, about 580°.

L, J. S. CINNAMALDEHYDE, cinnamic hyde, C.H. CH CH CHO. occurs number of esential oils, notably in cinnamon

and casus oils and forms an important article of commerce. It is extracted from oil of cunnamon by formation and subsequent decomposition of the crystallino sodium bisulphito compound (Penne, Ber 1884, 17, 2109) and from cassia oil by means of the double compound with barrion

aulphanilate (G P 124220).

Cinnamakieliado is most conveniently pre-parai synthetically by allowing a mixture of 10 parts of benzahlehyde, 15 parts of seetaldehyde, 000 parts of water, and 10 parts of 10% codium hydroxide solution to stand at 30° for 8-10 days with frequent shaking, it is then isolated by ether extraction (Peine, Ber 1884, 17, 2117, cf. B P. 10003) Two further synthetic methods of preparation from a chloroally benzeno bave been described (Bert and Doner, Compt. rend. 1930, 191, 332; Bert and Annequin, abid. 1931, 192, 1315) which are capable of extension to substituted cinnamaldehydes. It is best purified by means of the double compound with nitrie acid (Dumas and Péligot, Ann. Chim 1834, [u], 57, 322, Pfeiffer, Annalen, 1910, 376, 298)

It is a colourless oil with a pleasant emnamonlike odour; mp. -75°; bp. 252° (decomp), 130°/20 mm.; do 1 0497; np 1-6195. It alrould, theoretically, exist in two genmetrically isomeric forms, and Bourguel (Bull Soc. clum, 1929, [iv], 45, 1067) has described the pure ciscompound, bp 111°-113°/16 mm.; do 1 032; no 1.565, and its diethyl acetal. It is volatile in steam, practically insoluble in water and miscible in all proportions with alcohul. It may be characterised as its phenylhydratone, m p. 168°, semicarbacone, mp. 215°, and 24-disafro-phenylhydrazone, mp. 215°. It forms two oximes, syn., mp 130°, and antis, mp. 76°, the orientation of which has been established by

affords benzoic and acetic acida. Reduction recreatablised from water or alcohol.

cannamy! alcohol or y-pheny !propy! alcohol may be produced (Tuley and Adams, J. Amer, Chem, yields y-phenylpropyl alrohol (Shima, Mem, Coll Sci Kyoto, 1929, [A], 12, 69).

Cimnamaldehy de is estimated in essential oils volumetrically by means of hydroxylamine hydrochloride (Bennett and Donovan, Analyst, 1922, 47, 146) or sodium hisulphite (Gildemeister and lloffmann, "Die atherischen Ole," 1928, Vol I, p. 743) and gravimetrically with semi-oxamazido (Hanus, Z. Nahr. Genussm. 1903, 6, 817). See ALDEHYDES,

H. N. R.

CINNAMEIN v. BALSAMS, BAISAM OF PERU.

CINNAMIC ACIO. \$-Phenylacrylac acid, .H. CH CH CO.H. The separation of a CAH, CH CH CO.H. solid acid from old specimens of oil of cinnamon was first observed in 1780 by Trommsdorf, but the acid was regarded as benzoic seid until this error was corrected by Bizio in 1826. Cinnamic acid was first closely investigated by Dumas and Peligot (Annalen, 1835, 14, 50)

Cinnamic acid is very widely distributed in nature, the best known natural sources being bound atorax and tolu and Peru balsams. It is solated from storax by boiling with sodium hydroxide, followed by precipitation with b) droebloric acid and reery stallisation (Beil-tein

and Kuhlberg, Annalen, 1872, 163, 123; cf. von Miller, 181d 1877, 188, 196). It is formed by heating benzaldelijde with acetyl chloride at 120°-130° (Bertagnul, Annalen, 1856, 100, 128). This process was greatly modified and improved by Perkin (J C.S. 1877, 31, 388) who heated a mixture of benzaldehyde, acctic anhydride and anhydrous sodram acetate; this reaction, which is known as the Perkin reaction, is that generally employed in the laboratory. An improved procedure is described by Meyer and Beer (Monatch, 1913, 34, 651) and Bacharach and Brogan (J. Amer. Chem. Soc. 1928, 50, 3333) have shown that the addition of pyridine greatly increases the yield. The cinnamic acid need in the early processes for the manufacture of synthetic inchgo was prepared by this method, but Caro (G.P. 17487, 18232) showed that it could be prepared more cheaply by heating benzal chlorido with anhy drous sodium acetate; I part of benzal chloride and 2 to 3 parts of powdered fused sodium acetate are heated at 180°-200° for 10 to 20 hours in an autocline with mechanical aturing. The melt is mired Brady and Thomas (J.GS. 1922, 121, 2008).

In the air it is oxidized mainly to cumamic added the maintenance of the residual solution is acid (f. Pound and Pound, J. Physical Chem., differed hole, endified, and cooled. The pre1934, 283, 1015) whilst more drastic oxidation experience acid is filtered off and
filtered house and realized and Polarized cumamic acid is filtered off and

sodium hypochlorite or hypobromite (G.P. 21162; Ullmann, "Enzyklopadie der teehnischen Chemie," 1921, Vol. IX, p 613; Schorigin, Issaguliantz, and Gusseva, J. Gen. Chem. Russ 1931, 1, 506)

It may also be obtained by condensation of benraldeliyde with malonic acid in the presence of ammonia or certain amines (Knoevenagel, Ber 1898, 31, 2602; G.P. 97735, 164296).

Cinnamic acid forms colourless, monoclinic, prismatic crystals, in p 133°, bp 300°. It is soluble in 3,500 parts of water at 17°, in 42 parts of ethyl alcohol at 20°, in 17 parts of chloroform at 15°, and is volatile in steam. It may be characterised as its amide, m p. 148°, p bromophenacyl ester, in p 145°, and p-phenyl-phenacyl ester, in p 182°. It forms erystalline salts closely resembling those of benzoic neid. Oxidising agents convert it first into benzaldeliyde (distinction from benzoic acid) and afterwards into benzoic acid, the final products when potassium permanganato is used being plienyl-gly ceric acid, benzaldeliyde, and benzoie gly ceric acid, benzaldchyde, and benzoie and oxale acids (l'ittig and Ruer, Annalen, 1892, 288, 27). On fusion with caustic potash it yields a mixture of benzonte and acctate. It may be determined by direct titration, by absorption of bromino or by oxidation to benzoic acid (Nicholls, Analyst, 1928, 53,

It is now generally accepted that comamic acid exists in four forms (Stobbe, Ber. 1911, 44, 2739): (1) ordinary cumanuc acid, m p. 133°; (2) alloeumamic acid, m p. 68°; (3) and (4) two seeinnamic acids, m.p. 42° and 58°. Of these (1) is the trans. and and (2), (3), and (4) trimorphs of the cir aeid, (2) being stable and (3) and (1) labile (Bulmann, Ber. 1910, 43, 568). Meyer and Pukall (Z. physikal Chem. 1929, 145, 766). 360), however, conclude from work on the bronune addition of the various forms that the three last are truly isomeric and not merely polymorphic. This view is not accepted by Robinson and James (J.C.S. 1933, 1453) From physical data Disenblir and Hass (Z. physikal, Chem. 1935, 173, 249) conclude that the three cis neids are some special kind of rotational isomers; this view is contested by Meyer (ibid 1935, 174, 77).

In the polymerisation of einnamic acid the following equilibria are set np (Stobbe et al., Ber. 1919, 52, [B], 666, 1925, 58, [B], 1548, 2415, 2859; 1926, 59, [B], 2254):

Cinnancie and may be reduced to \$ phenylpropionic acid (hydrocinnamic acid) by a wide range of methods, among which may be men-importing special tioned cotalytic hydrogenation (Paul and pound perfumes Gerum, Ber. 1908, 41, 2277; Vavon, Compt.

Another process of technical importance rend. 1909, 149, 999; BJeschen, Rec. trav. consists in oxidising benzylideneacetone with chim. 1916, 35, 270, 277, 285), reduction with sodium amalgain (Alexejew and Erlenmeyer, Annalen, 1866, 137, 327) and with formic acid in the presence of palladium black (G P. 267306). It forms a dibromide, in p. 201°, with bromine in a variety of solvents (Sudborough and Thompson, JCS. 1903, 83, 669; Fittig and Binder, Annalen, 1879, 195, 140; Michael, J. pr. Chem. 1895, [11], 52, 292; Ber. 1901, 34, 3664), while addition of hydrogen bromide gives β bromo \beta-phenylpropionic acid (l'ittig and Binder, le) other halogen hydrides behaving similarly.

> Cinnamic acid condenses with hydrocarbons, e g. with benzene and sulphurie acid it forms phenylhydrindone (Liebermann and Hartmann, Ber. 1892, 25, 2124), and with phenols, eg. with plienol, it yields hydroxyphenylcou-

marın (ıdem., ıbıd. 957).

Lsters.-Stoermer and Sandow (Ber. 1920, 53, [B], 1283) have formulated the general rule that the esters of cis- forms of substituted einnamie acids havo lower boiling points than thoso derived from the corresponding trans-

Methyl cinnamate - Crystalline solid, in p

36°, bp 263°, 127°/10 mm.

Ethyl cinnamate is obtained directly by tho condensation of benzaldehyde and ethyl acetate by means of sodium (Claisen, Ber. 1890, 23, 977; Posner, J. pr. Chem. 1910, [11], 82, 435; B.P. 4969, 1890; G.P. 53671; Marvel and King, Organic Syntheses, 1929, 9, 38). It is a colourless liquid, bp. 271°, 144°/15 mm.

Substituted Connamic Acids .- 0-Nitrocinnamic acid was formerly of importance for the manufacture of synthetic indige. Nitration of the free acid (Müller, Annalen, 1882, 212, 126) or of the cthyl ester (Friedlander, "Fortschritte der Teerfarbenfabrikation," 1888, 1, 125) yields n mixture of the o- and p mitrocompounds which are separated by taking advantage of the insolubility of ethyl p-nitrocumamite in alcohol (Drewsen, Annalen, 1882, 212, 150). o Nitrocumamic acid exists in two forms, trans., m p 210°, and cis-, in.p 143°, the former being produced by nitration; similarly it is trans pnitrocinnamie neid, ni p. 286°, which is formed by nitration, the cis- acid niclting at 143° (see CARBOXYLIO ACIDS).

II. N. R.

CINNAMIC ALCOHOL, Cinnamyl alcohol C&H2 CH:CH CH2.OH

(AND ESTERS). Comanue alcohol 15 a synthetic perfume with a soft hyacinth odour, and is used as such and also in the form of its esters in modern perfumery. Pure cinnamic nleohol melts at about 35°, but commercial samples often contain traces of impurities which keep it in a liquid condition. B p. 257°-258°; d 36 1 0 302 ref index, about 1 580. Cinnainyl acctate, butyrate, and cinnamate which are used as synthetic pertinues have balsanic odours, and are of value in imparting special nondescript odours to com-E. J. P.

CINNAMON and CASSIA. In the where the outer surface of the bark stull penists United States the terms cinnamon and cassus The fracture as short and granular but fibrous are used almost indescriminately, sithough it as on the suche. The obour and taste are similar there recognised that, strictly speaking, they to belong to two enturyl separate trees. In Great states is also more stringent and sightly mucia-Britain they are treated as separate entities ginous. even in commerce, the price of cunnamon being even in commerce, the price of emnamon being Microscopic Examination of the two barks two or three times that of cassia. For shows many similarities. In true cinnamon the convenience the two spices are treated selecenchymatous cells form an uninterrupted together.

Cinnamon is the dried inner bark of Cinnamomum zeylanıcum Nees (Fam. Laurarem), a bark, corresponding to the portion used for tree which grows to a height of 20 to 30 first cumamon, the selerenchymatous cells do not and which is native to Ceylon In practive, form a continuous ring. These cork cells of the the young trees are pruned to produre a straight outer bark of cassia are much smaller than stem, which is then cut back and shoots are those of the inner portion. The sclerenchyallowed to grow out from the stump The matous rells of the phelloderm, or inner bark, branches are cut when they are between 1 and 2 em. in diameter , the bark is shit longitudinally on the inner walls, this thickening being more with two or more cuts and horizontally every conspicuous with cassia than with cinnamon 30 cm., and removed by copper or brass knives | The number of starch grains is about the same to prevent discoloration of the tannin. The in both barks, although cassia starch grains to prevent discoloration of the fannin The interpretability of the state of the sta with faint longitudinal strictions On the as flavouring materials and also for their carwith faint indictioning structure of the sea packeting materials and and our eurifuc, the longitudinal structures are minister preperties. Cases buck are used as more definite, and there are frequent sears and well as the barks for the distillation of the sometimes holes due to leave The bark is eventual oils, and the leaves of both trees are very brittle and breaka with a splintery fracture distilled for their cils, but these latter differ. The odour is fragrant and the taste swretish and considerably from the bark oils. warmly are matic.

occasional patches of a greenish colour showing separately in the following table

Microscopic Examination of the two barks ring, whereas in cassia there is an uninterrupted ring duo to the outer bark, but in the inner are slightly elongated tengentially, but thickened

Chemical Composition -Liverseege ("Adul-Cassia is the dried bark of Cinnamomum teration and Analysis of Foods and Drugs, Cassia Blume (Fam Lauracew), and less care is 1932, p. 427) Winton, Ogden and Mitchell (22nd taken with its cultration class with graums and Royal Boom. Exp. 180 and 1803, 2010, cunamon. The outer barks sparily removed by the contract of the contract

		Clas	FE04	Caesia				
Author	Liverscege	Winton, etc	Attagon	Bichardson	Uinton, etc	Arragon,	Richardson	
No of samples	approx. 15	6	2	2	20	2	4	
Moisture .	89-105	7-79-10 48	82-114	54-73	65-110	10 5-11 0	93-174	
Ash	39-6-0	4 2-6 0 0 02-0-13	35-43	34-15	3 0-5 2 0 03-2 42	19-29	25-82	
Cold alcohol	7-6-13 0	100-136			46-167			
Volatile oil .	05-11	07-16	25-28	0-8-11	0 9-5-1	26 1-3-1-5	05-35	
Fixed oil Starch and reducing	1-1-3 3	1-35-1 68	05-12	16-17	1-3-4-1	1-3-1-6	0.1-24	
substances Crude fibre		16 6-22 0 31-4-38 5	193-226 31-4-317	25 0-33 1	16 9-32 0 17 0-28 8	31·7-33 9 25 2-26 0	14 3-26	
Protein		3 2-4-1	34-43	30-38	33-54	31-34	26-45	

183 CITRALS.

oil, and this is probably determined most satisfactorily by the method of Cocking and Middleton (Quart. J. Pharm. 1935, 8, 435). The powdered bark is mixed with brine, distilled, and the vapours passed through the top of a condenser into a graduated tube, into which a small quantity of turpentine has been previously distilled to make the mixture lighter than water, the condensed water itself being returned to the distillation flask. An air inlet and outlet is provided by a side tube between the bottom of the condenser and the graduated tube. Cinnamon bark contains from 0.5 to 1.0%, and cassia bark from 0.5 to 2.0% of volatile oil, as distilled in commerce.

Standards.—The "British Pharmacopæia"

requires that einnamon shall contain not more than 7% of ash, and acid insoluble ash not more than 2%.

Adulteration.-Cassia bark is often supplied instead of the true cinnamon, whilst other species of cinnamon and cinnamon from the true Cinnamomum zeylanicum, but grown wild in Scychelles, are passed off for either bark. T. McL.

CINNAMON BARK, ESSENTIAL The oil distilled from the bark of OIL OF. Cinnamomum zeylanicum Nees (Fam. Lauraceæ), a small tree indigenous to and cultivated in Ceylon. An inferior oil is produced in the Scychelles, where the tree is cultivated. The oil is distilled mainly from the chips and refuse bark after the collection of the cinnamon quills for the spice trade. Formerly a mixture of bark and leaves was used for distillation, but the oil thus obtained contains a considerable amount of engenol and is not suitable for use in medicine. The finest oil is produced in limited

quantities and is an expensive product.

Constituents.—The main constituent is cinnamaldehyde. Pineue, phellandrene, and cymene are present, and traces of methyl-n-amyl ketone, benzaldehyde, nonaldehyde, furfural, linalool,

and caryophyllene.

Characters.—A pale yellow oil becoming darker on keeping, sp.gr. 1.000 to 1.010, opt. rot. 0° to -1° at 20° , $n_{\rm D}^{20}$ 1.565 to 1.591. Soluble in i volumes of 70% alcohol. Cinnamaldehyde content, 50 to 65% ("British Pharmacopæia" standard), determined by the hydroxylamine method. English distilled oils usually comply with these requirements, but less fragrant oils containing up to 75% are found in commerce at lower prices. Inferior oils containing admixtures of artificial cinnamaldehyde and distilled cassia oil are offered on the English market. It should be noted that the cinnamon oil official in the "United States Pharmacopæia" is redistilled cassia oil (q.v.). The Seychelles oil of cinnamon bark contains only 20 to 35% of cinnamaldehyde and is not soluble in 70% alcohol. C. T. B.

OF. The oil distilled from the leaves of ESSENTIAL On. or). It is produced in large citral b, by the oxidation of geraniel (Zeitschel, quantities in Ceylon and also in the Scychelles. Ber. 1906, 39, 1787).

The principal constituent is the essential It is used in soap perfumery and constitutes a valuable source of eugenol.

Constituents.—Eugenol, 1-a-pinene, 1-phellandrene, dipentene, benzaldehyde, linalol, terpineol, borneol, geraniol, safrole, a and β -caryophyllene, benzyl benzoate, einnamaldehyde, einnamyl alcohol, and traces of other alcohols are present.

Characters.—Sp.gr. 1.040 to 1.060, opt. rot. -1° to 3° at 20°, n_D²⁰ 1.530-1.545, phenolic constituents 70-90%. Soluble in 3 volumes of 70% alcohol. The phenolic constituents are determined by adsorption in the cold with aqueous 5% solution of potassium hydroxide.

C. T. B. CINNAMON-STONE or HESSONITE. A gem-variety of garnet of a warm reddishbrown colour (hence the name) and transparent. It is essentially an orthosilicate of calcium and aluminium, Ca, Al, Si, O12, containing, however, isomorphous replacements of iron, manganese and magnesium. Brilliant crystals are found in veins in serpentine at Ala in Piedmont, but material of the best gem-quality is found as pebbles in the gem-gravels of Ceylon.

L. J. S. CINOBUFAGIN v. CARDIAC GLYCOSIDES,

TOAD POISONS

CITOBARYUM. Prepared barium sulphate for use as a barium meal (Merck, Darmstadt, Napp, London), B.P.C. 1934.

CITRALS, THE. Citral a. (Geranial.) CMe₂:CH·CH₂·CH₂·CMe || HC·CHO CH2:CMe·CH2·CH2·CH2·CMe

Citral b. (Neral.) CMe₂:CH·CH₂·CH₂·CMe OHC·CH CH₂:CMe·CH₂·CH₂·CH₂·CMe || OHC·CH

Citral is the most important natural acyclic aldehyde of the terpene series. It occurs very widely distributed in nature and it forms the main constituent of lemon grass oil (from Cymbopogon flexuosus Stapf). It finds wide technical application both directly in the perfumo industry and also indirectly for synthetic purposes. The natural aldehyde is a mixture of the four aldehydes represented by the formulæ given at the head of the section.

Citral a, C₁₀H₁₆O, b.p. 118-119°/20 mm., d²⁰ 0.8898, n²⁰ 1.4891, is most readily obtained from lemon grass oil, of which it forms approxi-mately 90% of the aldehyde content, by the CINNAMON LEAF, ESSENTIAL OIL preparation of its crystalline sodium bisulphite OF. The oil distilled from the leaves of compound (Tiemann, Ber. 1899, 32, 117; Cinnamonum replanicum Nees (Fam. Lauraceæ) Hibbert and Cannon, J. Amer. Chem. Soc. differs entirely from the oil obtained from 1924, 46, 119). It yields a semicarbazone, m.p. the bark of the same tree (r. Cinnamon Bark, 164°. It is obtained also, almost free from 184 CITRALS.

Citral b. b p. 117-118°/120 mm., d' 0-8888, | product of citral and aniline with concentrated no 1-4951, semicarbazone, m p. 171°, is separated sulphurie acid gives a mixture of the cyclofrom citral o by taking advantage of the more ready reaction of the latter with cyanoacetic acid (Tiemann, Ber. 1899, 32, 120) It can be obtained eyche aldehydes, also by the oxidation of nerol. Both aldehydes CMe. are, when freshly distilled, colourless liquids with a characteristic odour ; they oxidise rapidly

on exposure to the air becoming yellow in colour.
The structures assigned to the two aldebydes follow from their preparation by the oxidation of the primary alcohols geraniol and nerol On oxidation with silver oxide citral yields geranic acid (Semmler, Ber. 1890, 23, 2965, 3556), which is obtained also by the conversion of citral oxime into geramonitrile followed by hydrolysis (Tiemann and Semmler, Ber 1893, 26, 2717). On exidation with potassium permanganate followed by chromic acid citral yields a mixture of acctone, levulinic and oxalic acids (Tiemann and Semmler, Ber. 1893, 28, In agreement with these results Harries (Ber. 1903, 36, 1933, 1907, 40, 2823; Annalen, 1905, 343, 351; d. Grignard, Docuvro and Escontrou, Bull Soc chim. 1924, 35, 932) has found that the citral semicarbazones, yield on oxidation with ozone, formaldehyde, acctone, lævuhnaldebyde and probably glyoxal. Con firmation of the structure assigned to the citral semicarbazone is hydrolysed. On oxida-aldehydes is provided also by their degradation tion it gives a mixture of β cyclogeranic and firmation of the structure assigned to the by alkali to methylheptenone.

Citral has been prepaced synthetically by the distillation of a mixture of calcium geranate and calcium formate (Tiemann, Ber. 1898, 31, 828).

of a number of derivatives and since in esential whilst from citral by pseudosomo, by 146-oils entral predominates derivatives of this 1507/15 mm. d* 0-593, mp. 153446, mp. 148-144, is obtained [Into the semicarbasones referred to above. suitable derivatives are a-citrylidene cyanoscetic acid, mp. 122°, heritylidene cyanoacetic acid, mp. 94°, and a cityd β-naphthocinchonic acid, m.p. 198-200°.

On catalytic hydrogenation in the presence of colloidal palladium citral yields a mixture of citronellol and citronellal (Shita, Ber. 1909, 42. 1634); and with platinum black a mixture of 2 6-dimethyloctano and 2 6 dimethyloctanol is formed (Vavon, Ann. Chim. 1914, [ix]. 1, 169). whilst with platinic oxide Adams and Garvey (J. Amer. Chem. Soc. 1926, 48, 477) obtained geraniol, citrouellol, and 2 6-dimethyloctanol.

Citrsl shows a marked tendency to undergo cyclisation passing into peymene in the presence of mineral acids. If, however, eather of the 1 omeric citrylidene eyanoacetic acids 13 absorption maxima at 504 and 473 mm in carbon heated with duiste sulphure and a mixture of chasplake. The structure (1) was proposed for an Af p cyclocarlysidene expansance cash as the structure (1) was proposed for otherwise, which yields on hydrolysis with alkala Acta, 1937, 20, 621; this has been established as and p cyclocarlas (1) and (1)), the high cyclocarlas (1) and (1), the high cyclocarlas (1) and (1), the leng formed in the larger quantity (Strubel, 530, 201) who effected the degradation of CP. 183331. Theman, Ber. 1909, 33, 3703 (2), 201) who effected the degradation of capasantian to circum by heating in a shown that the treatment of the condensation bydroziek.

citrals. Merling (Ber. 1908, 41, 2064) has also described methods for the preparation of these

For the apparation of a-cyclocitral, by, 90-95°/20 mm., d15 0-925, semicarbatone, mp. 206°, advantage is taken of the fact that when a mixture of the two aldehydes is reduced with sodium amalgam B-cyclocitral is converted into high boiling products whilst a cyclocitral is not attacked. It yields on oxidation a cyclogeranic acid, mp 1035°, It condenses with acetone to yield a sonone. & Cyclocitral, bp 112-114°/29 mm , d1225 0 9586, m2 1 1970, semicarbazone, m.p., 166-167°, is obtained free from a-cyclocitral by distilling a mixture of the semicarhazones of the two aldehydes in steam in the presence of phthalic acid when only β-cyclogecome acids. It condenses with acctone to

give \$ ionone. Citral a condenses with acetone to yield a pseudosonone, hp. 159-165°/20 mm, do Citral can be characterised by the preparation 0 8954, np 15317, semicarbazone, mp 1420. mann, Ber. 1898, 31, 2318; 1899, 32, 827; 1900, 23, 882; Friedlander III, 689; GP. 122466, 127661, 130457; Hibbert and Cannon, J. Amer. Chem. Soc. 1924, 46, 119) Analogous compounds with other Letones have been prepared (G.P. 150771).

J. L. S. pigment, which CITRAURIN. This separates in red needles from aqueous methyl separates in red needed from aductors metal-alcobol, in p. 144-1457, is one of several raro-tenoids which pigment the ripe orange (Citrus ourontinm), others being hyptoxantian, rea-zantian, lutein, carotene, and violaxantian (Zechmeister and Tuzion, Ber. 1936, 69, [B]. 1879). Citraurin, a polyene aldehyde, exhibits absorption maxima at 486 and 457 mu in carbon disulphide, the oxime m p. 181-182°, possesses heated with ddute sulphure acid a mixture of disulphase The structure (1) was proposed for

Citraurin is also obtained by the partial oxidation of zeaxanthin with potassium permanganate (Karrer, Solmssen and Gugelmann, Helv. Chim. Acta, 1937, 20, 1020).

I. M. H., R. F. P., and F. S. S.

CITRIC ACID, C.H.O. Occurrence. This acid occurs in the free state in the juice of many acid fruits such as lemons, currants, cranberries, limes, etc. It is also found free or as salts in the seeds and juices of many flowers and plants. It is a constituent of wine, milk, cheese, and animal tissues and fluids.

Synthesis.—The acid can be prepared from sym-dichloroacctone by treatment with hydrocyanic acid and hydrochloric acid, whereby dichloroacetonic acid is formed. This is then neutralised with sodium carbonate and heated with potassium eyanide. The resulting solution of dicyanoacetonic acid is saturated with hydro-genehloride and heated on a water bath for 15 hours. The citrio acid formed can be separated as calcium citrate by neutralising with milk of lime (Grimaux and Adam, Compt. rend. 1880, 90, 1252). It has been prepared by the action of potassium eyanide and hydrochlorio acid on ethyl acctonedicarboxylate (Dunschmann and von Peelmann, Annalen, 1891, 261, 162). Ethyl eitrate has been obtained in small quantities by heating together ethyl bromoacetate and cthyl oxalylacetato in the presence of zine (Lawrence, Chem. Soc. Proc. 1897, 13, 65).

Physical Properties .- Citric acid crystallises from cold aqueous solutions with 1 mol. of water. The erystals belong to the orthorhombic system (Groth, Chemische Krystallographie, 1910, 3, 475), and have a sp.gr. of 1.553. They are stable in air of ordinary humidity but lose water rapidly in dry air (Marchand, J. pr. Chem. 1842, [i], 23, 01). On warming slowly they soften at about 70°-75° with loss of water and finally melt completely between 135° and 152° (Buchner and Witter, Ber. 1892, 25, 1160). If the crystals are rapidly licated they melt at about 100°, then solidify as they become anhydrous, and melt sharply at 153° (Meyer, Ber. 1903, 36, 3599). The acid is obtained anhydrous by cooling its hot, concentrated solutions (Buchner and Witter. Le.; Meyer, Le.). The mean transition point at which the anhydrous and hydroated forms may reparato from solution is 36.3°±0.15° (Bennett and Yuill, J.C.S. 1935, 130). The latter workers givo a crystallographic description of the anhydrous form of citric acid which crystallises in the heleledral class of the monoclinic system. The melting-point of the anhydrous crystals is given as 156-157° and the density as dis (vac.) 1.665. Citric neid is optically inactive. dissociation constant of the first hydrogen atom is 8.2×10^{-4} at 18° (Kolthoff and Bosch, Rec. trav. chim. 1928, 47, 558) and the values for the record and third hydrogen atoms are given as 1.77×10⁻⁵ and 3.9×10⁻⁷ respectively. The ordinary crystallised acid dissolves in about half its weight of boiling water. Aqueous solutions have, according to Gerlach (Z. anal. Chem. 1869, 8, 295), the sp.gr. at 15° given in the table of lead of next column. in the table at head of next column,

	1	1	
Acid %	Sp.gr.	Acid %	Sp.gr.
1	1.0037	34	1.1422
$\overline{2}$	1.0074	35	1.1467
3	1.0111	36	1.1515
4	1.0149	37	1.1564
5	1.0186	38	1.1612
6	1.0227	39	1.1661
7	1.0268	40	1.1709
8	1.0309	41	1.1756
9	1.0350	42	1.1814
10	1.0392	43	1.1851
11	1-0431	44	1.1899
12	1.0470	45	1.1947
13	1.0509	46	1.1998
14	1.0549	47	1.2050
15	1.0588	48	1.2103
16	1.0632	49	1.2153
17	1.0675	50	1.2204
18	1.0718	51	1.2257
19	1.0762	52	1.2307
20	1.0805	53	1.2359
21	1.0848	54	1.2410
22	1.0889	55	1.2462
23	1.0930	56	1.2514
24	1.0972	57	1.2572
25	1.1014	58	1.2627
26	1.1060	59	1.2683
27	1.1106	60	1.2738
28	1.1152	61	1.2794
29	1-1198	62	1.2849
30	1.1244	63	1.2904
31	1.1288	64	1.2960
32	1.1333	65	1.3015
33	1.1378	66	1.3071
- 1			

and moderately so in ether. At 15° the solubility of the crystallised acid in 100 parts of alcohol is as follows: in 80% alcohol 87 parts of the acid; in 90% alcohol 52.85 parts; in 100% alcohol 75.90 parts (Schiff, Annalen, 1860, 113, 190; Bourgoin, Bull. Soc. chim. 1878, [ii], 29,

100 parts of anhydrous other dissolve 9.12 parts of the crystallised acid (Lippmann, Ber.

1879, 12, 1650).

Chemical Properties.-When heated to 175° citrie acid is partially converted into aconitic acid (Crasso, Annalen, 1840, 34, 56), the remainder losing water to form earbon dioxido and acctonedicar boxylic acid which immediately breaks down to carbon dioxido and acetono. Beyond 175° an oily distillate appears which yields crystals of ilaconic acid. At still higher temperatures citraconic anhydride is formed as an uncrystallisable oil (Anschutz, Ber. 1880, 18, 1541). When eitric acid is digested with slightly fuming sulphuric acid (von Pechmann, Ber. 1884, 17, 2542) or oxidised with potassium permanganate solution at 35° (Deniges, Compt. rend. 1900, 130, 32) acetonedicarboxylic acid is formed. At higher temperatures potassium permanganate solution gives oxalic acid (Phipson, J.C.S. 1862, 15, 142). By fusion with potassium hydroxide oxalic and acetic acids are formed A 25% solution boils at 101-8°, a 50% solution (Liebig, Annalen, 1838, 26, 158). Citrio acid at 105 8°. The acid is very soluble in alcohol is fermented by numerous moulds, yeasts and

saits of the aikaine earths are only aparingly soluble. The calcium sait, Ca₂(C₄H₂O_{7), 4H₂O, is less soluble in hot than in cold water and can} be dried at 100° without loss of water of crystallishtion. If the moisture content exceeds. 10-12% calcium citrate is hable to decompose when stored at room temperatures. When precipitated in a crystalline state its solibility in water is 1 in 1180 at 14°, and 1 in 1730 When theewn down in an amorat 90-100°. phons condition the solubility is greater, 1 m 707 at 18°, and 1 in 1,123 at 100° (Warington, JCS 1875, 28, 939) Citric send readily forms stabla doubla citrates and hence the hydeexides of many metals are not precipitated by alkalis in the presence of citric acid. Fernic ammonium citrate, which is used for medicinal purposes, is prepared by evaporating a solution of ferric citrate in ammonia on glass plates, the red compounds contain 20 5 to 22 5% Fe, and the green 14 to 16a' Fe

Acyl Derivatives and Esters .- The acelul derivative of cities anhydride is prepared by warming the acid with acetyl chloride on a water bath, dissolving it in lukewarm water and keeping in racuo over conventrated sulphurie The ethereal solution is removed, evaporated acid. The colourless syrup soon deposits shining crystals of acetoestric acid which are very solublo in water and melt at 135°-140° (Easterfield and Sell, J C S. 1892, 61, 1005) Hethylenecitric acid is prepared by heating citric acid with four times its weight of polymerie formaldehydo at 140'-160°, and allowing the melt to ervatallise Steinberg, Chem Zentr 1902, 1, 299) The pure crystals melt at 208°. Trimethyl citrate is obtained by bubbling hydrogen chloride into a mixture of anhydrous citric soid and absolute methyl alcohol for 2 hours, then refluxing for 6 hours, and finally crystallising from water and puryfying by distillation. It melts at 75.5° and boils at 98°/0.2 mm. (Donaldson, McCleary and Degering, J. Amer. Chem. Soc. 1934, 56, 459) In the same paperan account is given of the preparation of propyl, butyl, and amyldimethyl Asia Minor. Since the development of the curales by the alcoholysis of trimethyl curate industrial fermentation processes an increasing in the presence of the corresponding alcohol and quantity has been manufactured by these p-toluenesulphonie acid as a estalyst

Detection .- Pure citro acid is easily recog nised by the action of heat. It evolves an acrid odour and yields an oily distillate which forms crystals of staconic acid on rooling (Stevens, Ind. Eng Chem 1924, 16, 155). A solution of calcium chloride added to a neutral solution of are used for the preparation of juice and cutrie acid produces no precipitate in the rolld, essential oil. The fruits acc cut in half and the the calcium entrate. Ca₁(P₁(P₂)P₃)P₄(P₁), between the contains the essential oil. Its first account of solution on beiling Yarnous suggested methods for the detection of either seed in mutuure, depend on the insolubility of some of its mutuure, depend on the insolubility of some of its method is that such as the selection selection of either and present again before discarding and present again before discarding.

beterns (Kayser, Chem. Zentr. 1896, 1, 583; Compt. rend. 1900, 130, 32), gives acctone on Bruve. J. Biol. Chem. 1934, 197, 1191. Souling (Täufel and Schoierer, Z. Unters. Salits—Chris each is tribase and forms three classes of salits. The salits with the alkaline careful soluble in water; the neutral server soluble of the alkaline careful servers o a method using eitric dehydeegenase from eucumber seeds and tha methylene blue technique. It is difficult, however, to obtain concurdant results by any of these methods. Precipitation of the metallic salts fails to distinguish eitric acid unless it can be obtained fairly pure and in comparatively large quantity. Owing to the instability of acetonedicarboxviic acid even at room temperatures, methods depending on the formation of this compound are liable to give erratic results. Thunberg's method is only suitable for very small quantities of citrie acid and requires apecial technique. The various methods have been reviewed by Elsdon and Lees (Analyst, 1933, 58, 328) who consider that those depending on the formation of pentahromoaretone are tho most satisfactory, They have found that a preliminary extraction with ether increases the sensitivity of Kunz's pentabromoscetons test (Analyst, 1916, 41, 378) and recommend the following procedure; 200 g of the substance to be tested are treated with 50 e e of ether in a atoppered flask for 20 hours. to dryness, and the residue dissolved in 10 cc of water. This solution is treated with 1 cc. of sulphure acid(1:1), 0 3 ee of 37 5% potassum hromado solution, and 1 ec. of 5% potassum permanganate solution. The muxture is heated for 5 minutes at 45°. Any trace of manganese dioxide is removed by the addition of ferrous aulphate solution containing sulphure acid. If, after this treatment, any precipitate is present, citric acid is indicated. Under these conditions 01 g, of citro acid can be detected even in the presence of 100 g of tartane acid and 100 g, of sucrose

MANUFACTURE .-- For many years the main source of citric acid was the citrus fruits, chiefly lemons, pineapples, bergamots and limes, grown in Italy, Sicily, West Indies, California and Assa Minor. Since the development of the

methods Lemon, Bergamot and Lime Juices -Lemons are the principal source of citrus juice used for preparing citrie acid and they are obtained chiefly from Italy and Sicily. The best and ripest fruits are chosen for export and the rest in mixtures, depend on the insolubility of some of pulp is often fetched once of trein white. The state white the selection said (Kind, Analyst, 1900, 25, 40), the barram said [Brocelsmit, S.C.I. 1920, 29, 1733], and the lead said (Freenman and Grünhut, Analyst, 1913, 28, 55). Others depend upon the forms a properties of the state of the sta

used at once for the preparation of the acid. The juice is, however, usually concentrated until its sp.gr. is 1.24 and exported as such, in order to avoid the tariff imposed by many countries on the manufactured acid. Many manufacturers prefer to prepare the calcium salt, which is less citric acid, citrates and certain other acids, such as malic, aconitic, formic, acetic, propionic acids, and their salts are usually present in citrus inices. The acidity does not, therefore, give the true content of citric acid. Citric acid in such liquors is estimated by a method described later, depending on the separation of insoluble enleium eitrate. About 8% of the total seid in concentrated Sicilian lemon juice is not precipitable as calcium citrate. Con-centrated bergamot juice has a similar sp.gr. to lemon juice, but a lower acidity, usually about 32% $C_0H_0Q_1H_2Q$. About 13% of the total organic acid is not precipitable as calcium citrate. Concentrated lime juice has a sp.gr. of about 1.32 and an acidity of about 56% $C_cH_nO_7,H_2O$. About 10 to 14% is not precipitable as the calcium salt. The following table due to Grosjean (J.C.S. 1883, 43, 333) gives analyses of commercial samples of concentrated femon and other juices:

		as eitric	Acid (reckoned as eitric acid) in oz. per galion.				
	Density	Free acld,	Total acid precipi- table.	precipi- table to 100 of free acid.			
Lemon Juice: Average of 65							
samples	1.241	62.1	61.6	99-2			
Sample A .	1.240	65.8	59.7	90.7			
Sample B .	1.235	64.0	55.7	85.8			
Bergamot Juice							
Highest	1.235	47.0	48.5	101-4			
Lowest	1.235	52.3	49.9	95.4			
Lime Juice:	1						
Sample A .	1.326	108-3	99-8	92.2			
Sample B	1.205	59.2	53.9	91-1			
	t		1				

Citric acid is usually prepared by way of the calcium salt. The juice, if concentrated, is diluted with water and run into a vat fitted with a steam heater and stirrer. The temperature is raised to 60°, and sufficient hydrated lime of high nurity is added to neutralise 90% of the total acid estimated as citric acid. Chalk is then added to neutralise the remaining 10% of the total acid, and an excess of 7 kg. of chalk for every 3,700 litres of juice is also added. When the operation is completed a sample should no longer effervesce when more chalk is added. The liquid shows a slight acidity of about 0-15 to 0.20% CaH, O., H.O. This acidity remains however great the excess of chalk. When the neutralisation is finished the mixture is boiled about 90-95% of the iron and nickel present. for 5 minutes and the precipitated calcium The temperature is kept at 75° during the

containing about 4% of citric acid, and it can be | citrate washed with boiling water until the wash water has only a faint pale straw colour. It is not advisable to neutralise completely with lime, as dark-coloured impurities are thrown down which are afterwards difficult to remove. Both the lime and the chalk should be as pure as possible, since alumina, magnesia, iron and costly to pack and transport. Besides free phosphates interfere with the precipitation of citric acid, and cause appreciable losses of the acid. At a later stage in the process a washing of ealcium sulphate takes place. This wash is used as a medium in which to suspend the calcium citrate. This mixture is slowly run into a vat containing sufficient sulphuric acid to decompose the citrate and to leave an excess of not more than 0.2% sulphuric acid. The mixture is kept agitated and the temperaturo raised to 55-57°. The temperature should at no time exceed 60°. The acidity is roughly gauged by adding 5 e.c. of 45% calcium chloride solution to an equal volume of filtered liquor. After placing on a steam bath for 5 minutes a faint precipitate of calcium sulphate should be noticeable. The acid liquor is drawn off from the precipitate of calcium sulphate which is then washed free from acid. The wash is used as mentioned above and the acid liquor is concentrated in open pans until it has a density of about 20°-25°Bé., and is then filtered from the calcium sulphate which separates during the concentration. The concentration is then completed in vacuum pans until the density is about 37°-38°Bé. The liquor is run into a vat and kept agitated while cooling. By this process, known as granulation, a licavy crop of small crystals of citric acid is obtained. operation is a deliente one and requires to be done slowly. If the concentration is taken too far the result is a viscous liquid which will not granulate satisfactorily. The lumps of crystalline citric acid are allowed to drain before they undergo a light wash in a centrifuge. The mother liquor and the washings are concentrated again and a second erop of granulated material obtained. The process may be repeated a third time. When the liquor is too impure for further crystallisation it is known as "old liquor" and is returned to the neutralising tanks where it is treated like fresh juice.

The granulated citric acid is dissolved in water at 75° to give a solution of density 28°Bé., measured hot. A sample of the solution is filtered and a determination made of the charcoal required to decolourise the solution and of the quantity of calcium ferrocyanide to precipitate any iron and nickel present. determination is earried out as follows: to several 25 e.e. portions of the clear filtered liquid varying amounts of Ca₂Fe(CN)_{6.12H₂O} are added together with 0.5 g. of decolouring charcoal, and the mixtures heated on a steam bath for 10 minutes. The solutions are filtered and a drop of 1% Ca2Fe(CN)8,12H2O added to each. The sample which shows the least colour but still has a very faint blue, is used as the basis for calculating the amount of ferroeyanide to be added to the bulk. Usually enough ferrocyanide is added to precipitate

purification process. The filtered Equor is maintained if occasional growth cycles are introevaporated at low temperature until the density duced. This is accomplished by sub-culturing reaches 35°-36°Bd., measured at 50°. About on to small quantities of solution similar to that 2 to 3 hours before this point is reached a sample used in the technical fermentation and next on to is withdrawn and tested for free sulphume acid, fruits rich in vitamins and containing organic which, if present in large excess may appear in sends, e.g. oranges, lemons, tomators, etc. the finished crystals. For this test a sample of Kresling and Shtern (Proc. Inst. Sci. Res. Foed the filtered liquor is mixed with an equal volume Ind. Leningrad, 1935, (I), 3, 25) maintain of 45% calcium chloride solution and heated on a water bath. If more than a shight precipitate is formed a sufficient quantity of lime is added to reduce the excess to about 02% H,SO, If lead hard vessels have been used in the process appreciable amounts of lead may be present and can be removed at this stage together with any copper, tin and antimony. This is done by adding sulphuretted hydrogen water. By the end of the operation the sulphides will be precipitated and the exress of enlphuretted hydrogen boiled off,

The liquor is filtered clear and run into the crystallising vessels If large crystals are required the liquor is kept perfectly still for about 3 to 5 days. When small crystals are required the liquor is kept in constant motion. After crystallisation is complete the honor is drawn off and conventrated again. The process of crystallisation may be repeated as long as a satisfactory product results. The crystals are drained, washed in a rentrifuge, and dried at room temperature, or in a low-temperature

atore, before packing. Manufacture by Mould Fermentation --Wehmer (JSC I. 1894, 13, 275; BP. 5620, 1893) first found that citric acid could be obtained from augnr solutions by means of certain moulds. No auccessful industrial process was developed, however, until Thom and Curre action of many of these metals, Steinberg G. (J. Agric. Res. 1916, 7, 1) showed that strains a controversy about the similar of the mould Aperpillus may produce that was for the controvers of the mould Aperpillus may produce the trong time cortice and than Wehmer's original all necessary for normal growth and that these Citromyces. In a subsequent publication Curre december connot replace each other. To (J. Biol. Chem. 1917, 31, 15) laid the foundation for the commercial fermentation process by a and a strict control must be exercised on

There are hundreds of stroins within the species tion in their mochemical characteristics Different strains show great variations in their capacity for producing citric acid (Bernhauer, Biochem. Z. 1928, 197, 278). Those are selected quantities of suitable medium. Protod'yakonov and Kresling (Proc. Inst. Sci. Res. Food 130). The stock cultures must be sub-cultured by wt. calculated on the augar provided at definite intervals. Unless precantions are taken the acid producing capacity of the mould when the citric acid fermentation has been is hable to deteriorate. Stark (B.P. 302335) attempted on a commercial scale. Even if claims that the acid producing capacity of the the mechanic sternheed before inoculation it is

activity by occasionally sub culturing several generations of the mould on media containmg 0 025 to 0 05 N-hydrochloric seid. In order to prevent unsuspected contamination of the stock cultures a method of hactenological purification must be employed at definite intervals. Sucrose, invert sugar, glucose, maitose, and other sugars are suitable substrates for the fermentation. The best yields of citre for the fermentation. The best yields of citric acid are obtained, however, with sucrose. Currie considered the following medium to be most favourable for citrie acid fermentation with Aspergillus niger:

g per litre Sucrose 123-150 NH, NO, . 20-25 KH.PO. MgSO.7H.O . 0 75-1 0 . 0 20-0 23 HČI 4-5 re of N/5-solution so that the pn is 34-35.

The presence of metallic ions in the solution exercises a powerful influence on the growth of the mould and the production of and Stein berg (Amer. J Bot. 1919, 6, 330) gives a list of 17 elements, mainly heavy metals, which have been found to accelerate the development of Aspergillus. Although there has been considerable controversy about the atimulating mould is very sensitive to traces of these metals careful study of the optimum conditions for the their presence. Thus Smith (J.S.C.I. 1936, 55, formation of citric acid by Aspergillus nager 217T) at stea that the addition of 0.0001% of manganese to a culture solution rentaining Aspergillus niger, differing but slightly in their small amounts of iron, copper and zinc caused morphological details, but showing wide varia- large increase in growth of the myrelium of Aspergillus niger. Individual atrains show considerable differences as regards the optimum salt conventration for citric acid formation Vaniev (Brochem. Z. 1935, 278, 226) has which give the highest yields of the acid and do shown that the different strains of mould which not produce other acids when grown on small produce citric acid react differently from the addition of traces of zine sulphate. The optimum nutrient salt concentration has to be Ind. Leningrad, 1935, (I) 3, 3) state that the found for each particular strain used, and for strains which give low yields form notched, the type of vessel employed as the medium darkly pigmented conidia and irregular-shaped which gives the highest yields with small scale heads, while those giving high yields form fermentations is not necessarily the best when smoother heads. The latter are also more used on the large scale. Working with glass flasks active in synthesising starch and form less fat Bernhauer, Duda and Siebenäuger (Biochem Z. in the mycelium (Kresling, ibid. 1935, (1), 3, 1931, 230, 475) obtained yields as high as 76 3%

Various difficulties have been encountered atock cultures on gelatine or agar media can be difficult to prevent infection by bacteris, with the citric acid fermentation. Certain organisms have a particularly unfavourable effect on Appergillus niger. Yuill (Biochem. J. 1934, 28, 222) describes a Penicillium which contaminates Aspergillus niger cultures. Fernbach and Yuill (B.P. 266415) add sufficient hydrochloric acid to the medium to bring the p_{11} to about 1-8 at which concentration the selected Aspergillus niger can grow to the virtual exclusion of invading organisms so that heat sterilisation can be omitted. A better yield of citric acid results when the inversion of sucrose takes place graduelly and under the influence of the mould. The diminished hydrolysis of the sucrose and the reduced risk of metallic contamination of the medium from the reaction vessel are stressed in the above patent as advantages of the omission of heat sterilisation.

The fermentation is usually carried out in large, shallow vessels at a temperature of about 30° to 32°. The spores of the selected organism begin to germinate about 5 hours after they have been seeded on the surface of the medium The mycelium develops deep folds and wrinkles which present a very large surface to the culture solution. Citric acid rapidly accumulates as soon as the mycelium has developed. Tho whole of the solution is constantly brought into contact with the mould by stirring or other means and controlled quantities of air are admitted to the cultures. After 8 to 12 days the solution is run off, and the mould pressed and leached. The liquors are then treated as above for the preparation of citric acid from citrus juices. Wells and Herrick (Ind. Eng. Chem. 1938, 30, 255) report a yield of 63% by wt. on the sugar taken.

Whilst the highest yields are obtained using pure sucrose, eitric acid can be obtained from a cheap source of raw material such as molasses. the yield being about 19 per cent. Cabn (Ind. Eng. Chem. 1935, 27, 201) carries out the fermentation with Aspergillus niger on shallow layers of bagasse impregnated with a molasses mush. The layers of bagasso are about I to I in. thick and are supported on iron screens which form the bottoms of shallow trays. Aeration can thus take place both from above and below. The trays are protected from the acid by painting with a Bakelite varnish.

The fermentation occupies about 36 hours to 4 days, and sterilisation of the mash and its protection from infection from foreign organisms are stated to be unnecessary. At the end of the fermentation the citrie acid is leached from the bagasse carrier, which can be used over and over again.

METHODS OF ANALYSIS .- Commercial Citric Acid.—Accurately weigh out 3 g. of the sample, dissolve in about 100 c.e. of water, and titrate with N-sodium hydroxide using thymol blue 10 c.c. of concentrated sulphuric acid for I hour;

yeasts and other fungi which interfere scriously able substances). Dissolve 2 g. in 20 c.c. of a solution containing about 10% of NH, and I e.e. of 10% solntion of calcium chloride, and allow to stand for 24 hours; the solution remains clear (limit of oxalic acid). 2 g. in 40 e.e. of water and add 10 e.e. of dilute ammonia solution (as above) and 5 drops of 10% lead-free solution of sodium sulphide; the colour produced is at most only slightly deeper than that produced in a similar mixture containing in addition 1 c.e. of a 10% solution of potassium cyanide (limit of copper and iron). Dissolve 2.5 g. in water and add 1 c.c. of 10% barium chloride. Stir immediately with a glass rod and set aside for 5 minutes. The turbidity produced is not greater than that produced by adding 2.5 e.e. of N/100 sulphuric acid to a similar solution without the sample (limit of sulphates). Arsenie can be conveniently determined by the Gutzeit method and should not be present in quantities greater than I part per million. Lead is determined by dissolving in water 7 g. (primary solution) and 2 g. (auxiliary solution) of the sample; 10 e.e. of a solution of lead nitrate containing 0.001% of lead are added to the auxiliary solution. Each solution is made alkaline with a lead-free solution containing 10% of NH3, and then 1 c.c. of 10% lead-free potassium cyanide is added. If the solutions are turbid they are filtered. The two solutions are diluted to 50 c.c. and 2 drops of a 10% lead-free solution of sodium sulphide are added to cach and thoroughly mixed. If the the colour in the primary solution is greater than that in tho auxibary solution the sample contains more than the limit of 20 parts of lead per million. The proportion of lead in the sample can be determined by observing the quantity of the dilute solution of lead nitrate which must be added to the auxiliary solution in order that, after dilution to 50 c.e., there may be equal colours produced on the addition of 2 drops of the sodium sulphide solution. On incincration the sample should not leave more than 0.05% of residue.

Citrus Juices and Acid Liquors .- The acidity is determined by titration using thymol blue or phenolphthalein as indicator. Combined organic acids are estimated by neutralising a measured quantity of the sample with sodium hydroxide, evaporating to dryness, and gently igniting. The ash is dissolved in a known quantity of standard acid, boiled and filtered. The excess of acid is determined by thration with alkali. The alkali added when neutralising the sample, subtracted from the alkalinity of the ash, gives the amount of alkali corresponding to the bases of the organic salts. From this figure the amount of combined organic acid can be calculated.

Citric acid is usually estimated by Warington's method (J.C.S. 1878, 28, 934) or a modification the "British Pharmacopoia," 1932, gives
the following tests for purity: heat 1 g. of the powdered sample in a boiling water bath with didited to 50 c.c. and heated to boiling in a salt or glycerin bath and sufficient calcium not more than a faint yellow colour is produced chloride solution added to precipitate all the (absence of tartaric acid and of readily earbonis organic acid present. The whole is boiled

The filtrate and washings ero concentrated to ebout 10-15 c c, and the solution neutralised with a drop of ammonia if it has become acid. The second precipitate thus obtained is collected, the filtrate being used to transfer the precipitate on to the paper, which is washed with hot water. In very eccurate estimations the concentration should be repeated and any further precipitate collected. The precipitates in their papers are then burnt at a low red heat and the neutralising power of the ash determined with standard acid end alkalı. Each c c. of N-acid corresponds to 0-070 g. of C2H2O2H2O. Sometimes the first precipitate only is collected and, the conditions being standardised, allowance is made for the solubility of the coleium citrate

Calcium Citrate -Excess of chalk or hime is estimated by dissolving 5 g of the sample in dilute standard hydrochloric acid in a covered beaker and then titrating with standard alkali Citrate is estimated by gently igniting 2 g of the sample end determining the alkalmity of By the esh with standard acid and elkah deducting from the alkalizity of the ash that due to the chalk previously determined, the amount of base which has to be calculated as citrate is found. If other organic saits are present the citrate is dissolved in dilute hydrochlene and and the citrate estimated by Warngton's method as in the case of citrus

Juices Mill, Wine, ele -Citrie acid occurs only in small quantities in these substances Milk contains about 1 to 4 g, and wine about 0 4 g. per litre The methods of estimetion are based on the semo principles as those mentioned above in the section on "detection." There are no very simple methods of estimation and many of those described in the literature are unreliable Lampitt end Rocke (Analyst, 1936, 61, 654) have examined e number of the methods and recommend a technique for estimating citrie acid with special reference to milk. Their method is carried out as follows. to 50 e c. of milk serum (Lampitt end Bogod, Chim. et Ind 1932, 28, 777), or other solution containing citric acid, are edded 10 c c. of sulphure acid (1:1), if not already added in the preparation of the solution, except for pure citrie acid and milk serum, 10 c c, of freshly prepared bromine water should also be added, and any precipitate formed from acctonedicarbexylic acid filtered off after standing for half an hour. Potassium permanganate solution is added drop by drop from e burette with constant shaking until a brown precipitate persists, 10 ce being usually required for 0 1g, of estricaerd and 25 ce for a milk serum. The mixture is allowed to stand at room temperature for I hour, further potassium permanganato being added if the brown precipitate disappears. Sufficient 20% ferrous sulphate in 100 sulphuric acid is then added slowly until a pale yellow solution containing a white precipitate is obtained, end the mixture is cooled in an ice chest overnight. The precipitate is removed by filtration and the cedrino. The cedro has a thick rud through a sintered glass crucible, the reaction and hard flesh, the cedrono has abundant edible Sash being washed out with the filtrate to flesh, and the cedrino is a small fruit.

for half an bour and the precipitate collected remove the last traces of precipitate, and on a small filter and washed with hot water. the washings passed through the crucible. The precipitate in the crucible is then washed with 10, 10, and 5 c c. of cold water. The crucible is dried to constant weight in a vacuum desices. tor. The precipitate is dissolved out of the crucible with industrial spirit, followed by 20, 10, and 10 c c. portions of ether. The crouble is again dried in the vacuum desiccator and weighed, the loss in weight being taken as penta bromoscetone,

Catric scal, $C_4H_8O_9 = 0.424 \left(w + \frac{0.005V}{100}\right)$

to represents the difference in weight of the crucible before and after treatment with industrial spirit and ether, V the original volume of filtrate from the reaction mixture less the total volume of washings. Results show that citrie seid up to 011 g may be determined in the presence of lactose to within 2 mg.

B. T. CITRON. The fruit of Citrus Medica L. Originally of oriental origin, citrons ere now grown in Southern Europe and in California and Florida. The fleshy-welled fruit is not edible in the raw state but is well known as candied estron " The rind of the estma melon (Cstrullus vulgaris), a variety of the common weter-melon, is also prepared in a candled form as a substitute for the trus halved and fermented in brine for 40-50 days. during which the bitter flevour is removed and the tissue softened. The salt is then washed away and the peel steeped in a series of syrups of increasing sugar content until the maximum adsorption of sugar has taken place. Sucross cyrups tend to give a hard pecl, and an admixture of 35% of fructose is common. The composition of fresh and condied peel is given by Fellers and Smith (J. Agric Res 1930, 53, 859):

Fresh Candled 888 188 Ash 0-5 0.44 Protein . 0 15 0-16 0.33 Ether extract 0.321 37 Crude fibro. 1 09 43 O i Total sugars (as invert) 1 55 788 Total carbobydrates 94 0.09 0.06 CaO P,O, 0.05 0.04 0 11 0 02 KO. Fe, Cu, 1 traces

5 Consisting of 28 1% glucose and 14 9% sucrose A. G. Pa.

OIL CITRON. ESSENTIAL This oil, also known as oil of cedrat, is expressed in Sicily from the peel of the fruit of the citron, Catrus medica Linn (Fam. Rutacen). Three varieties are known as the cedro, the cedrone

C. T. B.

and a stearoptene identical with citrapteno

found in oil of lemon.

Characters.—A pale yellow oil, sp.gr. 0.850 to 0.852, opt. rot. at $20^{\circ} + 80^{\circ} - 81^{\circ}$, $n_{\rm D}^{2\circ}$ 1.475. Citron oil of commerce is frequently a mixture of oil of lemon, oil of orange and oil of bergamot. C. T. B.

CITRONELLA, ESSENTIAL OIL OF. The oil obtained by distillation from the grass Cymbopogon Nardus Rendle, which is prown principally in Ceylon, Burma and the Straits Settlements. Two varieties of this grass are found in Ceylon. The best quality, known as Maha Pengiri or Winter's grass, yields an oil containing a high percentage of acetylisable constituents, 80-90%, and is found in Burma and the Straits Settlements. The acetylisable constituents consist of geraniol and citronellal. The commoner variety, known as Lena Batu, yields the ordinary Ceylon oil, which contains 55 to 65% of acetylisable constituents, chiefly geraniol. Two or three harvests a year can be obtained under favourable conditions.

Constituents, -Geraniol, citronellal, camphene, depentene, limonene, borneol, nerol, farnesol, methyl heptenone, and an aliphatic sesquiterpene. Java oil contains, in addition, dicitronelloxide, methyl-eugenol and citronellic acid.

Characters.—Ceylon oil has sp.gr. 0.897 to 0.912, opt. rot. at 20° -6° to -14°, $n_{\rm D}^{20}$ 1.479 to 1.485. Total acctylisable constituents 57 to 62%. Soluble in 10 volumes of 80% alcohol. The best oils (so-called plantation oils) are clearly soluble in 3 to 4 volumes of 80% alcohol and remain clear with 10 volumes. Inferior oils become turbid with 3 to 4 volumes of 80% alcohol, clearing somowhat with 10 volumes. Such oils may contain 10-15% of petroleum. The solubility test on which London contracts are based requires that the oil shall be soluble in 10 volumes of 80% alcohol with only slight turbidity, and with separation of oily drops on standing 24 hours at 15.5°C.

Java oil lins sp.gr. 0.885 to 0.900, opt. rot. at $20^{\circ} - 2^{\circ}$ to -5° , $n_{\rm D}^{20}$ 1.468 to 1.473. Total acetylisable constituents not less than 85%, determined by the method of the "British Pharmacopoia" for the determination of free alcohols in essential oils, but using the formula

7.707n weight of acetylated oil-0 021n

-Percentage of acetylisable constituents,

where n=number of millilitres of N/2-alcoholic potnesium hydroxide solution consumed during the saponification of the acctylated oil.

Citronellal Content.—The Ceylon oil contains only 6-8% of citronellal and the determination is not usually carried out in commercial practice. For the dava oil, however, the determination of citronellal is of importance, and the method adopted is as follows:

Weigh accurately about 2 e.c. of the oil in a l stoppered tube and place in a freezing mixture. Wales, 1935, 68, 80). Add about 10 e.e. of a N-hydroxylamine hydro-

Composition.—The main constituents are chloride solution, prepared as below, and cooled limenenc, citral, of which 5 to 6% is present, to-10°, and titrate immediately with N-alcoholic KOH, adding the alkali slowly, a few drops at a time, taking great care to avoid going beyond tho orange colour. Continue the titration as long as a red colour develops, then allow to stand at laboratory temperature for one hour and complete the titration to the full yellow colour. Each c.c. of N-alcoholic potash is equivalent to 0.076 g. of citronellal.

N-Hydroxylamine Preparation | of chloride Solution .- Dissolve 6.95 g. of puro hydroxylamine hydrochloride in 95 c.c. of pure 90% (by volume) alcohol; add 10 drops of a 0.2% solution of pure methyl orange in 60% alcohol, and adjust to the full yellow colour of tho indicator with N/2-alcoholic potasb and make up to 100 e.c. with 90% alcohol.

The percentage usually present is 30 to 40. Both oils are used for soap scenting, and the Java oil is employed as a source of geraniol

and citronellal.

CITRONELLAL. The aldehyde, eitronellal, C10H18O, knownalso as rhodinal, although it is valued as a perfume, does not occur very widely distributed in nature. The dextrorotatory form, b.p. 205-206°, d^{20} 0.855, $n_{\rm D}$ 1.448, $[a]_{\rm D}$ +12.3°, is the main constituent of Ceylonese citronella oil from Cymbopogon Nardus, and it is found also in certain eucalyptus oils; the læveretatory aldehyde has been isolated from Javanese citronella oil and also from the oil from Pinus Jefferyi. For identification of the aldehyde tho semicarbazone, m.p. 85°-88°, may be used, but a more characteristic derivativo is eitronellyl-\betanaphthocinehonic acid, m.p. 225°, prepared by tho condensation of the aldehydo with β-naphthylamino and pyruvic acid (Deebner, Ber. 1894, 27, 2024). For the isolation of citronellal tho compound which it forms with sodium bi-

susceptible to the action of alkalis. The constitution of citronellal was established by the extended investigations of Harries and his collaborators (Ber. 1901, 34, 2981; 1908, 41, 2187; Annalen, 1915, 410, 8) on the oxidation of the aldehyde and its dimethyl acetal with ozone. They showed it to be a mixture of tho two aldehydes represented by (I) and (II), degradation products of both these forms being obtained as indicated in the schemo

sulphito may be utilised, but care must be taken

in its decomposition since the aldehyde is very

at the head of the next page.

Since citronellol (q.v.) gives on oxidation citronellal (Tiemann and Schmidt, Ber. 1897, 30, 33), the synthetic methods adopted for tho preparation of this alcohol apply also to the aldehyde. On catalytic hydrogenation citronellal yields the saturated aldehyde dihydrocitronellal (Paal, G.P. 298193) or the saturated alcohol (von Braun and Kaiser, Ber. 1923, 56, [B], 2271). On oxidation with silver oxide citronellic acid is obtained, the dl-form of which occurs in camphor oil (Rochussen, J. pr. Chem. 1922, [ii], 105, 124) and probably also in the oil from Calythrix Tetragona (Penfold, Ramage and Simonsen, J. Proc. Roy. Soc. New South

Citronellal condenses readily with acctone

OHC-CH,-CH,-CHMe-CH, CO,H

Мe Me,C:CH-CH, CH, CHMe CH, CHO Me,CO + CHO CH, CH, CHMe CH, CHO IT. HO, C.CH, CH, CHMe CH, CHO

in the presence of alkali to yield dikydro- (from Rosea damascena), geranium oil (from pseudoionone (III) and (IV) (Haarmann und Reimer, G.P. 75128; Rope, Ber. 1900, 33, 857) which on ring closure yields dikydroionone (V). This ketone has a refreshing odour and fieds

application in perfumery. Me,C CH CH, CH, CHMe CH, CH CH COMe III. MeCO HC CH

CH2-CH4 CH2 CH2-CH2 CHM4 CH2 CMCH COM4

Cyclic derivatives may be prepared also very readily from citronellal itself. As was shown by Tiemann and Schmidt (Ber. 1896, 29, 913; 1837, 30, 27) it yields on digestion with acetic anhydride isopulegol acetate, which can be hydrolysed to the alrohol (VI). The alcohol is formed also by the action of dilute sulphure and (Barbier and Léser, Compt. rend. 1897, 124, 1308; Horiuchi, Mem. Coll. Sci. Kyoto, 1028, 114, 171), formic and phosphoric acids (Prins, Chem. Weekblad, 1916, 14, 627). With aulnhuric acid the glycol (VII) is also formed. **CHMe CHMe**

СМе,∙ОН

manufacture of perfumes and occurs in a and Himmelmann (Ber. 1908, 44, 2187; Annalen, number of essential oils, such as rose oil 1915, 410, 8); authorquently Helferich and

Pelargonium odorantissimum) and in both Ceylon and Java extronells oils. It is identical with the alcohola known as rhodinol, reasol, roscol. The natural alcohols are optically active and the following constants have been observed: d-citronellol (from Java citronells oil), b.p. 103°/5 mm., d15 0-8604-0-8629, *1 1-4565-1-45791, [a]p +2 7°-2 3°; 1, bp 114-115°/12 mm., d29 0 856-0 862, n20 145609-14581, [a]_p -42°. The alcohol can be selentified most readily by the preparation of the allophanate, mp 105-106° (Granard and Docuvre, Compt. rend. 1928, 187, 270; Docurre,

Bull. Soc. chim. 1929, [iv], 45, 352)

The problem of the atructure of citronellel has formed the subject of a prolonged rontroversy now only of historio interest. Since it is formed by the reduction of citronellal (II or IIa) with sodium amalgam (Dodge, Amer. Chem. J. 1889, 11, 465; Tiemann and Schmidt, Ber. 1896, 29, 903) it must be a primary alcohol, and this is confirmed by its oxidation to extremellal with chromic acid, when at the same time citronellie acid (III and Illa) and sepulegol are formed (Tiemann and Schmidt, Ber. 1897, 30, 32). On oxidation with potassium permanganate followed by chromic acid, acetone, formic, and β methyladipic acids (IV) are obtained. The intermediate stages of the ovidation are shown in the scheme given at the head of the next page (Kots and Steche, J. pt.

Chem. 1924, [n] 107, 197). There can be no doubt that extremellol is mixture of the two alcohols represented by (1) and (In) the actual percentage composition of the mixture being dependent upon the source or prior treatment of the alcohol. In a careful series of experimenta Grignaed and Doeuvre (I e) have shown the alcohol from four different sources, (a) from Java extrenella oil, (b) from the reduction of citronellal, (c) from eitronellal acetate, and (d) synthetic, from methylheptenyl bromide, to be a maxture in which the alrohol (1) predominates, the percentage varying from 81 to 72%. These values are based upon the yields of J. I. S. actions and formaldehyde when circumdial conduct upon the years are to some upon the years are to conduct upon the years are to conduct upon the years are to conduct with ozone. The oxidation of circumdial conduction in the pello with zone was first studied by Harries

that the hydroxyaldchyde,

OHC·CH,·CH2·CHMe·CH2·CH2OH,

is formed also.

Citronellol has been synthesised in a number of ways: (a) by the reduction of ethyl geranate or ethyl eitronellate with sodium and alcohol (Bonvenult, Bull. Soc. chim. 1900, [iii], 23, 458; Compt. rend. 1904, 138, 1609; Barbier and Locquin, Compt. rend. 1913, 157, 1114); (b) by the reduction of eitronellal, and (c) by its conversion through its oxime into eitronellic acid with subsequent reduction of the ester of this acid; (d) from methylheptenyl bromido (Grignard and Doeuvre, l.c.); (e) by the reduction of geraniol with aluminium amalgam which is said to be used technically (Ullmann, Encyclopadie der Technischer Chemic, 9, 572). On entalytie hydrogenation eitronellol yields the enturated alcohol (Paul and Amberger, G.P. 316919: Haller and Martine, Compt. rend. 1905, 140, 1303). J. L. S.

Citronellol CITRONELLYL ESTERS. (q.r.) forms a number of esters valuable as perfumes. They may be described as having floral and fruity odours, with, in general, rose and bergamot predominating. The following are tho principal of these esters, with their approximate characters: acetate, b.p. 120°/15 mm., sp.gr. 0 901, np 1.4480; butyrate, b.p. 135°/5 mm., sp.gr. 0.891, n_D²⁰ 1.4490; propionate, sp.gr. 0.895, n_D^{20} 1.4455. Citronellyl formate, benzoate, cianamate, and valerianato are also prepared and used as perfumes to a small extent. E. J. P.

CITRULLINE, C. H13O3N3, an amino-ncid found in water-melon Citrullus vulgaris by Wada (Proc. Imp. Acad. Tokyo, 1930, 6, 15; Biochem. Z. 1930, 224, 420). It is formed by the action of putrefactive bacteria on arginine (q.r.) (Ackermann, Z. physiol. Chem. 1931, 203, 66), and yields ornithine on alkaline hydrolysis (Wada, Proc. Imp. Acad. Tokyo, 1972) 1932, 8, 367), and when perfused through the dog's stomach produces both arginine and ornithine. Citrulline has m.p. 220°-222°, is optically inactive and forms a violet copper salt, decomp. 257°-258°. The constitution c-amino-δ-carbamidovaleric acid,

NH2·CO·NH·[CH2]3·CH(NH2)·COOH, Vor. 111,-13

$$\begin{array}{c} \text{CH}_2\text{:CMe-CH}_2\text{-CH}_2\text{-CH}_2\text{-CHme-CH}_2\text{-CO}_2\text{H} \\ & IIIa. & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & \\ & & & & & & \\ & & & & \\$$

Sparmberg (Ber. 1931, 64, [B], 104) have shown | has been confirmed by synthesis from arginine by way of ornithine, dibenzoylornithine, δ-amino a-benzamidovalerie acid and 8-carbamido-abenzamidovalerie neid (Wada, l.c.).

> CIVET. Civet is the most obnoxious of the animal perfume materials, having a revolting odonr. When very highly diluted, however, its odour becomes pleasant, and its fixative properties in perfumes are very pronounced. It is a secretion from the perincal glands forming a deep pouch in the posterior part of the abdomen of the civet eat. The functions of the civet pouch are not definitely known, but it is probable that the repellent odour is a means of protection, or possibly its functions are those of sexual attraction. Practically the whole of tho world's supply comes from Abyssinia. Tho cat which yields most of the civet is Viverra civetta, a wild animal which becomes fairly tame on keeping in captivity, except when the eivet is being extracted. The cats are kept in long cages in which they cannot turn round, and their legs are secured and the civet extracted from the pouch with a small horn spoon. The secretion is greater if the animals are teased and irritated for a short time. There are other species of civet cats, but they are not of commercial importance. Civet is a substance of the consistency of honey, of a yellowish to brown colour, with a powerful ficeal odour. It is packed in bullocks' horns secured with a leather cap. The horns contain from 25 to 60 oz. Owing to its high price it is frequently adulterated. A genuine eivet should have the following characters:

Loss at 100°—not exceeding 25%. Mineral matter-not exceeding 2%. Acctone extract—65 to 80%. Alcohol extract—15 to 65%. Ether extract-11 to 24%. Chloroform extract—not exceeding 6%.

The acetone, alcohol, ether, and chloroform extracts should have the following acid and ester values:

		Acid value,	Ester value.
Acctone extract		85-115	25-50
Alcohol extract .		110-150	20-53
Ether extract .		8-26	55-90
Chloroform extract		6-10	75-170

The acetone extract should not melt completely below 40° and the suponification value CIVET.

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should be from 130 to 155. The principal given by Schimmel & Co. (Report, 1921, 60) as adulterants of civet are petroleum jelly, lanolme, showing the characters of a number of adulter butter and banana pulp. The following table is ated samples of civet:

CIVET (ADULTERATED).

_																
No			lextract		Ether extract.			L.	CH	Chloroform extract,			1 Kest-	Water,	Ash.	Mode of
30	*	AΥ	EV	s v	•	7.1	EΥ	sv	%	AV	EV.	s v.	due,	*	*	adulteration,
1	28 4	122 3	387	161 0	25 2	67	23 0	34 7	0:2	1-	I –	Ī-	16 9	293	12	Vaseline, organic
2	30 6	119-3	312	150 5	23-2	10 \$	33 9	19-1	\$0	8.	122 9	120 3	15 4	228	10	Vascline, organic substance, water
3 4 5	31 6 43 8 29 8 26 9	127 8 120 9 101 0 120 8	33 2 17 9 16 8 50 0	161 0 139 8 117 8 170 2	23 0 23 5 34 5 34 5	21 1 13 2 8 4 6 1	34 4	87 5 47 6 33 9 32 5	13		97 7 —	Ξ	68 68 168	87 71 23 284	1 04	Tale Vascline Vascline Vascline, organic
7	28 3	130 €	396	170-8	23 2	182	40 6	5S 8	0 5	-	-	-	176	30 4	-	Fubstance, water Vascune, organic substance, water
8	298	122 2	32 0	154 2	25 8	9-8	\$5.8	15 6	23	12 2	97 4	109 6	150	26 1	12	aneline, organic
10	11 5	119-1	36 9	156 0	١			840	3 4 0 5	=	=	=	5 8 20 2	25 3	15	Vascline, organic aubstance, water
11	35 3 33 3	1158	35 9	154 7	12 8 13 4	26.2	63.6	91.8	02	=	=	=	45 0 44 0	6 6 6 2	300	Tale Organic sub- stance
13	310	-	-	-	10 8	-	- 1		02	-		-	48 7	63	-	Organic sub-
14	33 5 37 6	129 7 119 9	27-2 53 6	157 9 173 5	11 0 19 6	39.6 30.1	63 7 71 4	89 1 100 0	05	=	=	=	48 1 19 3	22 0	42 2	Tale Sascline, organk substance, water

61, 901 , 1903, 62, 871; 1904, 65, 168), Burgess contain bydrated aluminium silicates. According (Analyst, 1903, 28, 101); Charabot and Hebert to J. W. Mellor ("Comprehensive Treatise on (Bull Soc. chim 1910, (iv), 8, 687), Schimmel Inorganic Chemistry," London, 1925, VI, 468 & Co (Reports, 1921, 60, 1924, 99, 1930, 110); there are three recognised hypotheses on the Niviere (Bull Soe chim 1920, (iv), 27, 794); and Bennett and Sesber (Perf & Essent Oil Rec 1929, 20, 14) Civet contains scatole, indole, ethylamine, propylamine, some free acids not identified, and a characteristic ketone which was described by Sack (Chem -Ztg 1915, 39, 538) under the name zibetone, but which is now usually known as civetone, melting at 31°.

Circtone has the formula C11H30O. It has

been completely synthesised by Ruzicka (Helv. Chim. Acts, 1926, 9, 230) and shown to be a 17-carbon ring compound Its constitution is as follows.

E. J. P.

CLARIFOIL. Cellulose acetate transparent tassuc

CLARIT. A German schwated earth. CLAUSTHALITE, a mineral of the formula

PbSe.

CLAVICEPSIN, C12H31O1a,2Aq, mp. 91°, and 198° (anhydrous), is found in Secule cornutum; it is not hydrolysed by emulsin, but gives 2 molecules of glucose and I molecule E. F. A.

"clay" denotes certain earthy materials which the different natural agents and the opportunited possess the property of plasticity and which which have occurred in geological time for such when heated to redness or higher become hard, processes as the leaching out or infiltration of

The following publications should be con Clays are the weathered products of silicate sulted E J Parry (Chemist and Druggist, 1902, rocks, especially those of the felspathic type, and conversion of felapathic cocks into clays. These are (1) the action of surface waters charged with carbon dioxide; (2) the action of water draining from peat bogs containing organic scids in solution; (3) the action of volcanic gases and vapours, that is, pneumatolytic action. In the normal weathering of granitic rocks, water, carbon dioxide and oxygen are the chief agents causing the breakdown. Ferrous compounds present in the rock are oxidised to hydrated ferme oxide, some of which may remain The alkah and sikaline earth compounds formed are removed in solution, while the quartz is comparstively unaffected. Thus, as the result of this weathering, by drated aluminium silicate, hydrated ferric oxide, quartz, together with proportion of felspathic or micaceous material, remain. If the water attacking the granitie rocks contains organic acids, a c. bumic acids in solution, the action is more severe. In some cases the seventy of the action has resulted in the breakdown of the hydrated sluminium sdicate, hydrated alumina remaining. Such an action is known as laterisation. Certain important beds of highly sluminous earths, such as themselves extremely diverse in character and CLAY (Fr. Argile; Ger. Thon). The term frequently very impure, owing to the action of

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seas, the admixture with mineral or organic impurities, sedimentation, the consolidation by pressure or earth movements. The china clays of Cornwall have probably been formed by pneumatolytic action. In support of this it is System considerable deposits of a series of pointed out that (1) the clay is often found at considerable depths in the body of the rock, (2) minerals containing boron and fluorine are

often found associated with these clays.

The terms "primary clay" and "residual clay" are used for those clays which are found at the place of their formation. China clay is an example of this type. In general, primary clays are comparatively free from impurities other than quartz and undecomposed felspars. These impurities are removed by washing from commercial china clay. A "secondary clay" or "sedimentary clay" is one which has been transported, usually by water, and redeposited. Such transportation and redeposition may have taken place many times. The sedimentary clays obviously fall into the geological classification of sedimentary rocks.

In Great Britain, in the Carboniferous System, there occur clays of considerable economic importance. Thus, in the Middle and Lower Coal Measures fireclays, which form the basis particularly Stourbridge, North Staffordshire, illustrate this diversity of composition:

soluble salts, the classification according to Yorkshire (Halifax, Leeds, Sheffield, etc.), particle size in streams, rivers, lakes and Durham, and Lanarkshire. Besides the manusers, the admixture with mineral or organic facture of firebricks, clays belonging to this class are used in the manufacture of drainpipes, glazed bricks, and sanitary fireclay ware. In the Upper Coal Measures of the Carboniferous clays known as Etruria marls are found in North and South Staffordshire and the Ruabon district of North Wales. These clays, which may be fired "red" or "blue" according to the methods of firing adopted, are used extensively in the production of roofing tiles. The Keuper marls (clays) of the Triassie; the Lias clays, the Oxford clays and the Kinimeridge clays of the Jurassie; the Weald clay and the Gault of the Cretaceous, and certain Tertiary clays, are all used in the manufacture of building bricks. Ball clays, which are characterised by high plasticity and bonding power, are fine-grained, sedimentary clays of the Tertiary group. They are found at Newton Abbot (Devon) and Poole (Dorset), and together with china clay are used extensively in the pottery industry.

THE COMPOSITION OF CLAY.-Most clays are composite bodies containing, in varying amounts, (1) hydrated aluminium silicates, (2) quartz, (3) ferruginous compounds, including limonite, siderite, pyrites, and nontronite, (4) alkaline earth compounds, (5) felspathie and micaccons of the refractory materials industry, are found. substances, (6) carbonaccous matter. The These clays are obtained in many districts, following ultimate analyses of some clays will

TABLE 1 .- ANALYSES OF CLAYS.

	China clay (Cornwall).	Ball clay (Devon),	Fireclay (Stour- bridge).	Fireclay (Scotland).	Etruria mari (N. Statis.).	Triassle clay (Birming- ham).
SIO:	9,4 46.90 0.37 39.52 0.30 0.25 0.16 0.20 0.06 12.56	59.42 0.94 27.24 0.62 0.47 0.22 2.26 0.40 8.90	% 55:24 1:34 28:35 2:25 0:40 0:22 0:84 0:36 10:52	% 44·24 1·36 36·92 2·16 0·22 · 0·14 0·36 0·16 14·10	50·10 1·54 24·64 10·72 2·10 0·46 1·82 0·36 8·20	54 ^{'3} 2 0·78 20·14 6·32 5·71 3·21 1·40 0·32 8·02
	100.32	100.47	l 99·52	99.66	99-94	100-42
	Ca	lculated Pro	ximate Ana	lyses.		
" Clay substance "	98.1	60·9 21·0 16·0	67·6 19·5 7·2	90-90 0-50 3-10	<u>-</u>	

(The titanium, iron, calcium, and magnesium compounds are not included in the proximate analyses.)

Numerous attempts have been made to examine clays by the so-called method of rational analysis, the object being to determine by chemical means alone the amounts of "clay substance," quartz, and "felspathic matter" present in them. These efforts have not proved satisfactory. By making certain not proved satisfactory. By making certain the "clay substance" clays ubstance "clays ubstance" has a formula Al₂O₃·2SiO₂·2H₂O, and that

ie. that given by the formula

J. W. Mellor (Trans Ceram. Soc. 1917, 16, 73) showed that a crystalline mineral—kaolimite -with the above formula could be isolated from china clay. He used the term "clayste" for the supposed non-crystalline "clay substance, which formed by far the greater proportion of clays he investigated. More recently C S. Ross and P. F. Kerr (US Gool Survey, Prof. Paper 165E, 1931) concluded that the crystalline fraction of Laolin comprises at least three distinct minerals, kaolinite, nacrite, and diekite, all with the formula Al₂O₂ 2SlO₂ 2H₂O—thus explaining the anomalous results obtained by previous workers. They also emphasised the presence of the non-crystalline fraction. It is to contain kaolinite at all. accepted in these days that other hydrated accepted in these days that other syndrict simminum sulcates may be present in class There include helloyate (AI, O, 280, 4H, O), allophane (AI, O, 510, AH, O), montmorollonite (AI, O, 480, H, O+2H, O), bedelinte (AI, O, 480, H, O+2H, O), pyrophylate (AI, O, 480, H, O+2H, O). It is interesting to note that there is a complete isomorphous arrive ranging from beidellite to nontcomite

the latter being often found in clays. Further, the X ray spectra of montmorillonite, beidellite, the A ray spectra of monumental commonly present in bedeflite, elay a containing and propoliphic are practically scientical commonly present in bedeflite, elay a containing and proportion continuous as the precognised as important continuous as the factory and the facto bentonites which are highly colloidal aubstancea firing coloura. Montmorillonite exerts an action

R. E. Grim and R. H. Birsy (J. Amer Cernar Soc. 1936, 19, 307) has o investigated the mineral constitution of thirteen examic elays of widely different properties. They pead particular the lags and ball clay are kaolantic clays; different properties. They pead particular the lags bonding power of the latter results attention to the colloid portion, which was more from high subdivision of the particles repided. turn subdivided into (-1+0 1µ), (-0 1+0 00µ), with the presence of organic material and some and (1-06g) fractions. A centriture technique benefitie. The senette like marral, other a fit. II. Bray, R. E. Gram and P. F. Kerr, domainst constituent of shales, under similar beind evised for this purpose. The different values for the unfired preperties that isolated marrals when the different values for the unfired preperties that isolated has to contain a potable and its alumining a second contains potable and its alumining as the contains potable and as the contains p the petrographic microscope and chemical analysis of the hulk sample and all fractions. X-ray diffraction photographs were also obtained on the colleidal fractions. The nonclay minerals such as quartz, mica, and glauconits were availty found mainly in the non-colloid observed, a retardation in the heating curred or portion. The clay mineral stated above comparing the colloid fractions, and generally acceptation in the rate of rise of temperatures of clay attacked. Somewhat similar results have 1910, 9, 91; 1911, 10, 161; 1914, 13, 83) made 1936, 24, 117). It is interesting to record the indiamentally crystaline character of practically all the inorganic clay constituents of colloidal dimensions which these works. ite were usually found mainly in the non-colloid

of which appreximated to that of china clay | examined. One mineral occurring in the colloid fraction in most of the clays examined has not been identified so far. It is described as "senutelike "and is apparently the same as that named "glammerton" (mica clay) by K. Endell, U. Hefmann and E. Maegdefrau (Zement, 1935, 24, 625). Grim and Bray have tentatively assigned to it the formula

2K.O 3MO 8R.O. 24SIO. 12H.O.

while Endell and co-workers state that it loses water continuously on heating between 200° and 600°C. Reviewing the evidence which they have collected, Grim and Bray have concluded that kaolinite, beidellite, montmorillomite, the sericite like mineral, and halloysite are the most important clay minerals. According to them, many clays and shales do not appear

Different ceramic properties are necessarily associated with the presence of these minerals in natural clays, but the effects are difficult to evaluate because of the influence of particle airo as well as the apecific attributes of each mineral In a coughly qualitative manner certain influences have been assigned. Thus, drying shrinkage and bonding strength increase with the content of beidellite. A characteristic of this mineral is its high base exchange capacity, henre clays in which it is present in quantity will be highly sensitive to additions of electeolytes Sinco alkalis and iron are benconties which are nighty consistent allousteness in mice groups. Another product the definition of the continuous cont probably replaceable by iron, it is not refractory

and as often red burning. THE ACTION OF HEAT ON CLAYS .- The effect of heat on clays was early examined by H. L. Le Chatcher (Z physikal, Chem. 1887, 1, 396). He observed a retardation in the heating curve of crystalline kaohinte at about 550°C, and an colloidal dimensions which these workers Table 2 gives the thermal data for these re-

actions obtained by W. M. Cohn (J. Amer. Ceram. Soc. 1924, 7, 359, 474, 548; Ber. deut. Keram. Ges. 1923, 4, 55); A. E. MacGee (J. Amer. Ceram. Soc. 1926, 9, 206); H. E. Simpson (J. Amer. Ceram. Soc. 1927, 10, 897).

TABLE 2.

Substance.	FOLL	t nb- otion 1 g. 0°C.).	evol	eat ution 1 g.			
China clay Georgia kaolin Zettlitz kaolin Halle clay Fireclays	93 127 92 93 82	cals.	16 23 16 11 18	eals.	W. M. Colm A. E. MacGee W. M. Cohn W. M. Colm H. E. Simpson		

Quantitative values for the above reactions have also been obtained by L. Navias (J. Amer. Ceram. Soc. 1923, 6, 1268). In addition to the above thermal reactions S. Satoh (Sei. Rep. Tohoku, 1923, (3), 1, 157) concluded from a study of his heating curves that a very weak exothermic reaction takes place between 1,170° and 1,220°C.

The endothermie reaction observed with many elays in the region of 500°C, is associated with the evolution of the combined water or water of constitution. Varying results for the dehydratien temperature of clays have been obtained, owing probably to the influence of the rate of heating. Thus, H. S. Houldsworth and J. W. Cobb (Trans. Ceram. Soc. 1923, 22, 111) found that kaelin lost a small proportion of combined water on heating to 400°C. in about 7 days, but the main loss occurred between 420° and 450°C. It has been further indicated by R. Ricke (Sprechsnal, 1911, 44, 637) that knolin is not completely dehydrated at 600°C. C. S. Ross and P. F. Kerr (U.S. Geol. Survey, Prof. Paper, 165E, 1930) have indicated that pure knolinite loses the largest part of its total combined water at about 450°C., dickite at about 575°C., and nacrite at about 650°C. Some clay minerals such as montinorillenite lose a considerable amount of their water at temperatures of 140°C. This water is probably different ia nature from the water of constitution of kaolinite. Rehydration below 400°C, is considered to be due to the presence of colloidal constituents such as colloidal silica, colloidal alumina, or "allophanoid" substances. Ross and Kerr have suggested that the water removed on heating the beidellite and montmorillomite minerals is composed of " zeolitic " water, water of constitution, and adsorbed water. The chemical change which proceeds at about 500°C. also influences the combination of the alumina, tendering this constituent soluble in hydrochloric acid. J. W. Mellor and A. D. Holdcroft (Trans. Ceram. Soc. 1910, 9, 94) consider that these observations are best explained by assuming the complete breakdown of the clay molecule into free silica, free alumina, and water.

assume that the water of combination is set free, leaving Al₂O₃·2SlO₂—leverrierite (W. Vernadski, Trans. Ceram. Soc. 1923, 22, 398; and F. W. Clarke, Bull. U.S. Gool. Survey, 1895, 125, 32)—or that an aluminium silicate of the type Al₂O₃·SiO₂, together with free silien, is formed (J. Samoilov, Trans. Ceram. Soc. 1924, 23, 338); whilst J. M. Knoto (Trans. Amer. Ceram. Soc. 1910, 12, 350) suggested the formation of two compounds, Al2O3 SlO2 and Al2O3.3S1O2.

In the neighbourhood of 850° the alumina becomes much less soluble in hydrochloric acid. The chemical change occurring between 850° and 1,060° has been attributed by J. W. Mellor to the polymerisation and erystallisation of the free alumina. Numerous observations on the formation of a crystalline alumino-silicate in clay bodies at high temperatures have been made. J. W. Meller and A. Scott (Trans. Ceram. Soc. 1924, 23, 327), on firing mixtures of kaolin and alumina between 1,200° and 1,700°, and afterwards treating the material so formed with hydrofluoric acid, obtained an insoluble residue approximating to the composition of silhmanite (Al₂O₃·SiO₂) after the treatment at the lower temperatures, and mullito

(3Al₂O₃·2SiO₂)

after the treatment at the higher temperatures. The more recent work on the X-ray spectra of kaolin after heating to various temperatures has led to different conclusions. J. W. Mellor, W. Bragg and G. Shearer (Trans. Ceram. Soc. 1924, 23, 314) stato that when kaolinite is heated at about 550° the crystal pattern of kaolinite disappears and is not replaced by any new pattern, and thus the products of decomposition are devoid of crystallino structure. The results obtained by J. F. Hyslop (Trans. Ceram. Soc. 1925, 24, 402) are somewhat different. The pattern obtained from china clay heated at 450° for 5 hours showed no marked change in the position of the kaolinite lines as obtained with the raw clay, but showed a decreaso in intensity. After heating to 550°C., a faint pettern unlike that of alumina or any of the silien modifications appeared, becoming more definite at 600° and persisting at 850°C. This pattern was considered to be due to a compound of the formula Al₂O₃·2SIO₂. After heating to 950°C., the pattern disappeared and was replaced by that of mullite (3Al₂O₃·2SiO₂), which became more intense after heating to 1,050°C. With clays containing quartz, heating at the higher temperature eaused the transformation of some of the quartz to cristobalite. J. F. Hyslop and H. P. Rooksby (Trans. Ceram. Soc. 1928, 27, 93, 299) in a later investigation confirmed the above observations and in addition noted the appearance of a pattern, associated with erystalline alumina, after heating to 870°C. This phase disappeared after heating to 1,060°C. Many other investigations have indicated the appearance of mullite in clays fired to 1,050 C, or over. In most cases the presence of either cristobalite or tridynute-crystalline This view is supported by G. Tammann and W. Pape (Z. anorg. Chem. 1923, 127, 43). Some of which, after heating to very high Other views which have been suggested

tory materials, building bricks, and other silics and alumina and the disposition of the rerame products are impure; in lace, town so called clays contain less than 40% of "elay ferric oxide, and thereby produce a paier but substance." This variation in the constitution colour, at the same time conferring greater substance. The substance on the firms operation, stability on the free oxide. The alumns, which reramic products are impure; in fact, some so- alumina to enter into solid solution with the principle of which is to produce from a for any cause, may not have taken up the iron satisfactorily manufactured and dried clay shape a finished product possessing strength, shape, and such general or particular properties as fit it for its subsequent duties. Since hardly any two elays are the same in their behaviour towards firing, it must srem that the firing operation is not capable of much standardisation Nevertheless, in considering the industrial firing operation, three definite periods of importance stand out. These are (1) the watersmoking period, ranging in normal firing from ordinary temperatures to about 250°C. , (2) the oxidation period ranging from 500° to 850°C.; (3) the vitrification range from 800° to the finishing temperature (v A T Green, Trans. Cerani Soc 1927, 26, 110) The water-smoking period is concerned with the expulsion of the water from the setting of the dried goods in the kiln, for, even after drying, the waco may contain up to 5% of water. In a kiln, full of heavy elay ware weighing 100 tons, this means that up to 5 tons of water may require to be chminated The inadequate appreciation of the control of this stage of the firing may not only set up severe strums and ultimate weakness in the final product but may also cause the control of the subsequent heat work to be scriously impaired The oxidation period is mainly concerned with the elimination of carboneceous matter, the oxidation of iron compounds, the breakdown of carbonates, sulphides, and other compounds, together with the decomposition of the rlay molecule with the evolution of the water of constitution During this stage many pro-perties are determined in the firing of bracks made from fireday, carbonaccous matter and the iron compounds may either individually or together form "black cores" or "dis-coloured centres". The nature of the icon oxide which forms during firing depends upon the amount of oxygen available, the steam in the atmosphere and the rate of rise of temperature. and highly micaceous clays this range is very In a limited supply of oxygen or in an excess of small, being of the order of 50", but with steam, ferrous compounds will be formed or good fireelaya the range during which appreciable maintained, while under strongly oxidising effects are evident isoften from 1,280° to 1,400°C. conditions, during the oxidation period, ferrie Here, then, is the crux of successful heat work compounds will be formed The formation of for it is essential to produce such an amount of " black cores" is often due to a deficiency of air vitrified matter which gives atrength with resili during this stage, so that the carbonaceous matter of the clay is incompletely exidised, and the iron remains in the ferrous condition forming a fusible ferrous ailicate, which causes chemistry of arrested reactions an early vitrification in the interior of the pro-It is obvious, therefore, that the rate of rise of temperature during this period must be fixed to different temperatures angue guidant most carefully controlled and that the steam information concerning its behaviour during formed owing to the decomposition of the dely the virturication. Contraction data are to the control of the control of the decomposition of the dely the virturication. Contraction data are to the control of the delay of the de formed owing to the decomposition of the clay the vitrification. Contraction dats are very must be quickly chiminated, the kiln atmosphere useful in the manufacture of clay products thus having to be carefully regulated.

As already stated, most clays—other than coloured fireclay bricks. It is based on the china clay—used in the manufacture of refractions of the clay at 500°-600°C, into free into solution (e g the iron being present as FeO or Fe,O, owing to reducing conditions), is less hable to do so at temperatures over 800°C. On this basis a rational explanation of bricks with red cores and buff exteriors, and with haff cores and red exteriors, has been

given. The final period of firing is that associated with the vitrification of the product. Vitrification may be defined as the progressive partial fusion of the various mineral constituents of a clay, chiefly including the felspathic, micaccous, calcareous, and fereuginous substances, which is governed by, amongst other factors, the composition of the body as a whole, the size of the grain, and the desired texture of the ware. fact, the object of the heat work during this period is the production of a suitable amount of vitrified or glassy material, which will curround the remaining particles and fill the insterstices to an extent dependent on the peoperties desired in the finished product Naturally, then, vitrification, which is the partial fusion of certain of the constituents of the clay resulting in the coherence of the on mass, depends on both chemical and physical factors. Physically, the grain size has a very considerable influence, for the greater the surface per unit area, the greater the interaction taking place. Chemically, the nature and amount of the fluxes present in the day determine the rate and extent of the reactions possible. With fireclass the rate of vitrification is allow up to temperatures in the neighbourhood of 1,200°C. Many clays used in the manufacture of building bricks such as Oxford clay sor Triassie clays show rapid vitrification at temperatures ranging from 900° to 1020°C. The range of temperature during which the actions involved in vitrification take place to any appreciable extent varies very considerably. With calcareous ence without affecting the shape of the product. Thus, J. W. Mellor has pointed out that the chemistry of the firing of clay wares is the

The measurement of (1) the change in perouty and (2) the linear contraction of a clay when which must be made reasonably true to are and J. W. Mellor (Trans Ceram. Soc. 1933, 82, shape, F. R. Ennos and A. Srott (Mem. 403) has brought forward an explanation of Geol. Survey, Freelays, 1925, 28, 3) have given certain colour effects due to iron oxide in buff. the porosity-temperature curves of many

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groups depending on the rate of decrease of porosity with temperature. Such a classification gives some indication of the maturing conditions necessary to produce a satisfactory product. In certain instances the clays show a stendy decrease in porosity with temperature of firing, pointing to a prolonged vitrification, whereas with others there is obviously a temperature at which the elay starts to vitrify with great rapidity.

It has already been indicated that siliceous fireclays contain free quartz varying in grain size from sub-microscopic particles to grains of appreciable magnitude. After firing, much of this quartz may remain as such embedded in the matrix or supporting structure of the fired body. Depending on the duration of the fire, the size of the grain, and the nature of the matrix, some of the quartz may be transformed iato eristohalite. It does not follow, however, that a high silica content in the original elay will produce high quartz or cristobalite content ia the fired product. The quartz, at least some of it, may go into solution in the viscous flux during vitrification and remain in this state after cooling. When silien is thus dissolved, the characteristic reversible expansions associated with the presence of quartz or cristohalite

disappear. THE PLASTICITY OF CLAYS; SUSPENSIONS .-The plasticity of clay may be defined as that property which cambles it to change its shape under the action of a deforming stress without cracking and without any tendency to resume its original form on the removal of the stress. In certain theories plasticity has been attributed to (1) the size and shape of the particles; (2) the physical properties of the water film surrounding the clay particles; and (3) the colloidal contents of the clay. Clays differ widely in plasticity, but in general such clays as hall clays and bentonites possess the greatest plasticity. Certain fireclays, in their natural coadition, are shaly, and require fine grinding with water to develop plasticity. Some clays, e.g. certain aluminous fircelays and chian clays, show little plasticity, these being technically described as "lean" or "short" as opposed to ball clays which are termed "fat." The amount of water required to produce maximum workability varies widely with the clay, "short" clays, in general, requiring less water than "fat" clays. J. W. Mellor (Trans. Ceram. Soc. 1922, 21, 95) has shown that the water content required to produce maximum workability decreases with the pressure applied to the clay in working. In the plastic, stiffplastic, and semi-dry processes of manufacture this fact is utilised. A clay which requires, say, 26% of water for the plastic method may be successfully moulded under high pressure (200 kg. per sq. cm.) with a water content of, say, 6%. Although no satisfactory method of obtaining an absolute measure of plasticity exists, many empirical tests have been used. In works practice, plasticity is usually gauged by " feel,"

clays. They divided clays into a number of the measurement of the flow of clays through dies and tubes, and the measurement of the effects of tension and compression on clays containing controlled amounts of water. A highly plastic clay has, in many cases, a high tensile strength in the dried state. This fact is sometimes used as the basis of an indication of plasticity.

E. C. Bingham ("Fluidity and Plasticity," New York, 1922) has developed a formula for the plastic behaviour of solids in which the plastic flow is related to the applied stress, yield value and the viscosity coefficient. T. W. Talwalkar and C. W. Parmelee (J. Amer. Ceram. Soc. 1927, 10, 670) have studied the stress-strain relation of clays by means of the torsion testing of cylinders of clay and have found the Bingham equation satisfactory. In these investigations important technical characteristics of different clays have been indicated. E. O. Wilson (J. Amer. Ceram. Soc. 1936, 19, 115) has also investigated plasticity by means of a torsion machine giving stress-strain relationships. results indicate that plasticity is due to the presence of stable viscous water films held on the surface of the particles; these in turn depend on the shape and surface activity of the particles. Clays show the highest plasticity because in addition to a plate-like structure they have a high surface activity. The older plate-structure and the colloid theories are thus blended. R. E. Wilson and F. P. Hall (J. Amer. Ceram. Soc. 1922, 5, 916), G. W. Scott Blair and E. M. Crowther (J. Phys. Chem. 1929, 33, 321), G. W. Scott Blair (J. Rheology, 1930, 1, 127; Trans. Ceram. Soc. 1931, 30, 138) have reported valuable investigations which have a bearing on the fundamental conception of the plasticity of clays.

> The workability of a wet clay may be increased by storing in a cool, damp place for a period of time. This process, known as "souring," gives a more even distribution of water throughout the mass and makes the clay tougher. Souring may be hastened by the addition of small quantities of certain organic substances, such as guins, tannin and humus, which favour the growth of certain bacteria. D. P. Glick (J. Amer. Ceram. Soc. 1936, 19, 169) has studied the micro-organisms present in six types of elay. By comparing the properties of raw, aged and sterile clays he has shown that the presence of living organisms are contributory factors to the ageing of clays. Traces of some salts affect plasticity considerably, a small quantity of acid increasing the workability and strength, whereas the addition of alkali has an opposite effect. The plasticity and drying shrinkage may be decreased by the addition of non-plastic material, such as sand or fired clay material (grog), and by suitable blending of different substances bodies of varying degrees can be obtained.

Elutriation determinations show that 68-98% by weight of ball clays consists of particles smaller than 0-010 mm. diameter. H. G. Schurecht (J. Amer. Ceram. Soc. 1920, 3, 355) found that an English ball clay contained 81% a manual treatment of pulling, bending, twist- by weight of particles of size less than 0.0005 mm., ing and wedging being used. Laboratory tests whilst some English china clays contained only include the Vicat needle test (r. Vol. II, p. 144a), 5% of such particles. J. J. T. Schlasing (Compt.

1-4% of material of a colloidal nature from a certain clay, whilst J. W. Mellor (Trans. Ceram. Soc. 1922, 21, 95) removed only 0.5% from another. In general, clays in suspension have an acid reaction, the pn range according to F. P. Hall (J. Amer. Ceram. Soc. 1923, 6, 901) being Jan 10. After Cream Soc. 1933, 0, soap camp. 3 10.7-37. Methods for the more accurate measurement of the particle size of clays have been put forward by A. H. M. Andreasen (Ber. deut Keram. Ges. 1930, 11, 249, 678), E. T. Knapp (Ind. Eng. Chem. 1934, 6, 66), C. R. Amberg [J. Amer. Ceram. Soc. 1936, 19, 207). The particles in a clay slip are negatively charged and can be deposited on an anode immersed in the clay slip. On suspending a clay in distilled water, the coarse particles settle out rapidly and the clay matter remains suspended for a length of time, the suspension floor. In this method the moulded articles are containing particles smaller than 11 micron This auspension is typically colloidal, and exhibita the Brownian movement, Tyndall effect, and cataphoreus Additions of small quantities of seid to a clay slip cause flocculation; the acidity at which maximum flocculation occurs varies with different clays, according to Hall, from a PR of 27 to 40 At this point the negative charge on the clay has been neutralised Small additions of alkalis to a clay alip defloces late the clay, the particles remaining suspended for a longer time, and the viscosity of the slip being reduced. If alkali be added in excess the viscosity is increased and the clay becomes florculated. Thus, a casting slip, which ordinarily would be too thick to pour, can be successfully east by the addition of controlled quantities of acdium silicate and sodium titanium, colcium, magnesium and the alkali carbonate Protectivo colloids such as gelatin (sodium and potassium) oxides, together with the and gum arabic, when added to a clay slip, may retard flocculation. Clays show selective adretard flocculation. Clay a show eelective and Many other constituents, e.g. the oxides of scorption; R. F. Geller and D. R. Caldwell manganese and barum, sulphure and photos. Amer. Ceram Sec. 1921, 4, 463 found that phore enhy drides, carbon dioxide and carbon, kaolins could adsorb sodium hydroxide to the

such as barium chloride and copper sulphate, permissible error are usually taken to be in preference to the acid ions, whilst larger ±0.5%, though different analysts work to molecules of organic dyes, e.g. methylene blue, different limits. If the analysis falls sensibly are also adsorbed The mechanism of the are also adsorbed. The mechanism of the below 89.5% a qualitative examination must be reaction of clays slips to electrolytra has been made to identify the constituents which have discussed by C I Whitlatch (J. Amer Ceram | escaped determination in the normal course of Soc. 1931, 14, 154), F. L. Clark (Trans Ceram. Soc. 1933, 32, 1), I) L. Peek and D A McLean (Ind. Eng Chem 1934, 8, 85) and Z W. Wolkowa

(Kollad Z 1934, 67, 280).

THE DRIVE OF CLAY -The abrinkage of clay during drying from the plastic atstetakea place in two principal stages. During the first the change in volume is equal to the volume of water lost; at a certain mossture content the clay becomes what is technically tabout 3 to 4 hours is generally aufficiently known as "leather hard," and at this point It is not usual to determine the hydroscopic the clay particles are in contact and shrinkage practically ceases During the second stage, from "leather hard" to dry, there is a small residual shrinkage attributed to loss of colloidal the dry material is heated in a platinum crucible water and to closer packing of the clay grains The transition from the first to the second stage minutes over a good Meker burner, with the is not audden, there being an intermediate stage, which varies considerably with different clays. and contents are cooled in a draiceator, weighed,

Throughout the first stage, use saw constant conditions of temperature of a clay under constant conditions of temperature of a clay under constant for unit are of surface, for during this stage there is a comparatively free flow of water to the surface. When the "leather-hard" stage is reached the free flow of the water to the surface is prevented. with a consequent falling off in the rate of drying. The shrinkage of the surface of a block of clay relative to the interior causes tensional strains to be set up in this auperficial material. If drying is too rapid these strains become great enough to eause supture and cracking. Care has to be taken that the greatest safe rate of drying is not exceeded. This safe rate depends

on the texture of the clay or clay-ware One of the commonest methods of drying elay products in industry is by means of the hot placed directly on the heated floor of a large shed. The chamber diver consists of an enclosed space with means for heating and ventilating In the tunnel dryer the goods are carried, and dried by atenm pipes, hot gases, or other means All types of commercial dryers are subject to very wide variations in design to meet the requirements of different branches of the clay industry. II. II. Macey (Trans, Ceram. Soc 1934, 33, 92), T K. Sherwood and E. W. Cominge (Ind Eng Chem 1933, 25, 1, III. I34) end F. II. Norton (J. Amer Ceram Soc. 1933, 16, 86) have made useful contributions to the study of the drying of clays.

ANALYSIS OF CLAYS .- General -The constituents determined in a normal clay analysis are silica, the oxides of aluminium, imn,

loss on ignition. may have to be determined in special cases. The last two are included under the less on extent of 1 23% on the dry weight of clay. The last two are included under the less on Clays adsorb the basic ions of common salts, ignition in a typical analysis. The limits of

the quantitative work. Sampling -It is imperative to sample esrefully the finely ground meterial, otherwise the analysis will not be a representative one

and consequently of little value.

Determination of the Hygroscopic Mosture - About 5 g, of the finely ground material are dried in an air-bath or tolucne oven at 105° to 110°C until there is no further loss in weight (about 3 to 4 hours is generally aufficient). mossture as normally the analysis is made on the dued sample,

Determination of the Loss on Ignition .- I g of for 15 minutes over a small flame, then for 30 lid on for the last 10 minutes. The crucible

burner and the weight again checked after

Low results are usually due to the incomplete combustion of carbonaceous matter, whereas losses by spurting, produced by too rapid ignition (especially if carbonates be present),

generally account for high results.

Determination of the Silica .- The ignited material in the platinum crucible is intimately mixed with 10 to 15 g. of anhydrous sodium earbonate. (Some analysts prefer to weigh out a fresh gram of the dry un-ignited material.) The lid is placed on the crueible and the mixture gently heated over a Méker burner and finally fused at a bright red heat until the contents are in a state of quiet fusion, usually 15 to 30 minutes are required. The crucible is allowed to cool on a clean unglazed tile; it is then half-filled with water and earefully heated over the tip of a small flame whereby the cake can usually be detached en bloc from the crucible. The cake and washings from the crueible are placed in a dry 250 e.e. basin, glazed on the inside only; after adding about 100 e.c. of water the basin is covered with a clock glass and 25 to 30 e.c. of concentrated hydrochloric acid aro added from a pipette through the lip of the basin. When the first violent reaction is over the basin is warmed on a water bath until all action has ceased and the cake has disintegrated. Any drops on the underside of the clock glass are rinsed into the basin; the platinum crucible and lid are washed with hydrochloric acid and hot water and the washings also poured into the basin. cake is now crushed to powder with a small agate pestle and the solution evaporated to dryness on a water bath until the smell of hydrogen chloride is no longer perceptible. When crystallisation has started the semi-solid mass must be repeatedly broken up with the end of a glass rod. The basin and contents are now baked in an air oven at 120°C, for I hour. The mass is moistened with concentrated hydrochloric acid and 75 c.c. of hot water added. After a few minutes' digestion the liquid is filtered and the residue washed with hot water until free from chlorides. The filtrate is returned to the basin, again evaporated to dryness, baked, digested with hydrochloric acid and hot water and filtered as in the first instance. The wet filter papers are transferred to a weighed platinum erucible and carefully dried and charred without ignition over a burner with a mushroom head. The carbon is then burnt off slowly over a Bunsen burner and the crucible and contents finally ignited for 30 minutes over a Méker burner, with the lid in position for the last 10 minutes.

The crucible is weighed and the result entered as "weight of crucible plus silica and residue." The silica contains traces of the oxides of aluminium, iron and titanium. Accordingly, it is treated with two or three e.e. of dilute sulphurie acid to prevent the subsequent volatilisation of titanic fluoride at red heat. About 15 c.c. of hydrofluoric acid are added, a few drops at a time, and, after placing the crucible eccentrically on a sand bath, the solution is slowly evaporated to dryness. When all the hydrofluoric acid has been expelled, the sulphuric acid can be rapidly precipitation only is necessary.

again heated for 10 minutes over the Méker and safely volatilised by heating the rim of the crueible, carefully, with the naked flame. The outside of the crucible is wiped free from sand; the erneible is then ignited for 5 minutes and weighed when cool. The result is recorded as "weight of crucible plus residue." The difference between the two weighings gives tho amount of silica in the sample. Subsequently the ammonia precipitate is ignited in this cruciblo along with the silica residue.

Normally the weight of the silica residue does not exceed 10 mg.; a residue much in excess of this figure usually indicates the presence of barium sulphate or of relatively large quantities of titanic oxide in the material under analysis. In such cases the silica residue is fused with a small quantity of potassium pyrosulphate, the cake extracted and digested with dilute sulphurie acid until completely disintegrated. The solution is filtered; the residue, if any, washed free from sulphates, ignited and weighed as barium sulphate. The filtrate from

the barium sulphate is added to the main bulk of filtrate from the silica.

The Ammonia Precipitate.—The filtrato from the silica is heated to 80° to 90°C, and 2 to 3 g. of solid ammonium chloride are added to it, followed by a slight excess of concentrated ammonia, drop by drop; the solution is simultaneously stirred well and then filtered, after standing for 5 minutes. If the elay contains manganese, which will be indicated by the green colour of the cake after fusion with sodium carbonate, 2 to 3 c.e. of bromino water aro added to the solution before the two ammonia precipitations, otherwise the procedure is as described above. The precipitato is filtered and immediately washed four or five times with hot water; a hole is made in the apex of the filter paper and the precipitate washed back into the beaker from which it has just been filtered.1 The filter paper is then washed free from chlorides and kept for ignition. The washings are collected in the beaker containing the precipitate. The precipitate is redissolved in a slight excess of concentrated hydrochloric acid; the solution, after dilution, heated to 80° to 90°C, and again precipitated by adding excess of ammonia, drop by drop with constant stirring. The liquid is filtered into the beaker containing the first filtrate and the precipitate washed repeatedly with small quantities of a hot alkaline solution of ammonium nitrate (2 to 3 g. per litre, made just alkaline with ammonia) until free from chlorides, but on no account must the ammonia precipitate be allowed to run dry.

The filtrate is evaporated to about 75 e.c. and 2 to 3 c.c. of ammonia are added and the excess boiled off. The liquid is filtered and the precipitate washed free from chlorides with the annuonium nitrate solution. The filtrate is kept for the determination of lime and magnesia. The three filter papers used in the animonia precipitation are transferred to the crucible containing the silica residue and the papers

 $^{^{1}}$ When the ammonia precipitate does not exceed $5^{\circ}_{\rm c}$, only a slight excess of aminonia is added to the filtrate from the silica and the excess is boiled off. One

are slowly dried and charred over a mushroom hurger. The temperature is then raised until above mentioned dilutions, the percentage of all the carbon is burnt off and the crucible Fe₈O₂ is given by V/8, where V is the number of finally heated, with the hd on, over a Méker e.e. of diluted standard iron solution required to burner until its weight is constant.

After weighing, the ignited buides are very slowly dissolved in the crucible by fusion with 6 to 8 g. of pure fused potassium pyrosulphate.
The cale when cold is detached from the

cruesble and transferred to a porcelam basin and about 150 c.c. of water and 20 c c. of concentrated sulphure acid are added to it; the crucible is well rinsed out with hot water and the washings poured in the basin. The mixture is heated on a water bath until all is dissolved; the solution is then cooled and diluted to 250 c c. in a graduated flask to give the stock solution for the determination of iron and titanium oxides

Determination of Ferric Oxide-When the amount of ferric oxide does not exceed about to it from a hurette, with frequent surring, 5%, a colorimetric process is used, while for larger amounts the determination is made by the ordinary permanganate process. The colour of the sample in the crucible after the determina tion of the loss on ignition is a good criterion as TiO_s is 25/(10+V), where V is the number of to which process is the better. For the coloriec, of water added to the 10 cc. of diluted metric determination an aliquot portion of the stock colution, obtained from the pyrosulphate fusion, is diluted to 250 cc. in a graduated diluted to 100 c.c. the percentage is . flask to give the test solution 25 cc. are taken for a normal clay, or other materials, containing 1% or less of ferric oxide. 5 c.c. of the standard iron solution 1 are diluted to 100 e e in a graduated flask and a portion of this solution is transferred to a burette reading to 0.1 c c., while a similar burette is filled with distilled water. Two small test glasses, of pure white glass, are respectively filled with a mixture of (a) 10 cc. of the potassium thiocyanate solution with 10 cc. of a buffer solution of potash alum a and (b) 10 cc. of the potassium thiocyanate solution with 10 cc. of the test solution The diluted etandard iron solution is added from the burette to the poissh alum solution and an equivalent amount of distilled water from the other burette to mixture (6), stirring thoroughly after each addition, until the tints in the two test glasses are the same. The tints are viewed against a neutral background by light transmitted through the aides of the permanganate produced in the oxidation, the glasses. will have reached its maximum intensity. The

If more than 6 to 8 c.c. of the diluted iron solution are needed, errors are likely to result owing to the difficulty in judging the equality of tint in concentrated solutions of ferme thiocyanate. A weaker test solution must, therefore, be made by diluting 10 or 5 c c. of the atock solution to 250 c.c.

¹Slandard Iron Soluton—Dissolve 0 6048 g. of pure ammonium ferrie alum in water, add 5 e. o. of conventirate uniputure acid, and distret he solution to a limit of the conventirate uniputure acid, and district he solution and in the convention of the convention o trat solution on the tint of the ferric thior; anate,

Calculation,-Using 1 g. of material and the give a match in colour. If the 250 c c. of test solution contains v e c. of the stock solution, the

percentage of Fe,O, is 25 V/8 v.

Determination of Tilanic Oxide. -50 c c. of the atock solution, obtained from the pyrosulphate fusion, are pipetted into a 100 cc. graduated flask, 10 cc. of 20 vols. hydrogen peroxide (free from fluorides) are added, and the mixture made up to the mark. 5 c.c. of standard tstanium solution 1 and 10 c.c. of hydrogen perexide are similarly diluted to 100 c.c. One of the test glasses used in the iron determination is about half filled with the colution to be tested. 10 c c. of diinted standard titanium solution sre preetted into the other glass and water added until the tints of the two liquids, compared as indicated in the previous determination, match,

Calculation .- If 1 g. of material be taken, with the dilutions given above, the percentage of standard titanium solution to produce equality of tint. When v c c of the stock solution are

1250/(10+V)r.

Determination of Alumina .- The weight of the ammonia precipitate less the weights of the ferrio and titanic oxides and the filter ashes, need in the silica separation and the ammonia precipitations, gives the weight of alumins in

the sample. Determination of Managnese Oxide -The percentaga of manganous oxide, though rarely required in a clay analysis, can be estimated colorimetrically as follows.

A suitable quantity—say 5 c c —of a standard manganeso cointion is pipetted into a 100 c c. graduated flask and to it is added 10 cc. of a 0-20% solution of silver nitrata and 1 g of ammonium persulphata. The flash is heated on a water bath until a pink colour develops By the time the contents of the flask have cooled to atmospherio temperature the colour, due to solution is then diluted to 100 cc. A brown precipitate on oxidation with the persulphate abows that insufficient silver nitrate has been used. In this case the solution is reduced by adding a few e c. of sulphureus and and reoxidesed, after the addition of another 10 cc. of the silver nitrate solution.

The stock colution, resulting from the pyroaulphate fusion, generally contains traces of

Standard Telantum Solution -1 g of pure titanic 1 Standard Tutassum Salaton —1 g of pure United to fixed with 10 g of potassium prosubblet. The cold cake is extracted with water and 20 cut and the cold cake is extracted with water and 20 cut and

Hence 50 e.e. of it are boiled and the chlorides precipitated by the addition of silver nitrate. The solution is filtered and the precipitate washed, the filtrate and washings being collected

in a 100 e.e. graduated flask.

10 e.c. of 0.2% silver nitrate and 1 g. of ammonium persulphate are added to the filtrate and the mixture warmed, cooled and diluted to the mark, as in the preparation of the standard permanganate solution. Tho tints of standard and tests solutions are then matched exactly as described under the colorimetric determination of titanie oxide.

Calculation .- If the volume of water, taken to dilute the standard solution to the same depth of colour as the test solution, bo v c.e., then, with the dilutions given above, the percentage of manganous exide in the elay is $2\cdot 5/(10+v)$.

Determination of Lime.—The filtrate from the

ammonia precipitate is boiled for a few minutes with 1 to 3 g. of ammonium oxalate, and 5 to 10 e.c. of concentrated ammonia are added. After stirring well, the mixture is warmed on a water bath for 2 hours. The liquid is filtered, the precipitate washed two or three times and the filtrate reserved for the magnesia determination. A hole is made in the apex of the filter paper, the precipitate washed into the beaker from which it has been filtered, and redissolved by running a few e.e. of nitric acid over the filter paper into the beaker containing the precipitate. The filter paper, after washing, is rejected. The solution is boiled, again precipitated and warmed for 2 hours as before, after which it is filtered and the precipitate washed free from chlorides. The filtrate is added to that from the first precipitation. Tho precipitate is slowly charred and then ignited in a platinum crucible and finally heated over a Meker burner for 15 minutes, with the lid on the erneible for the last 5 minutes. The erneible, when cool, is rapidly weighed, again ignited for 5 minutes and the weight checked.

The percentage of lime is calculated from the weight of the residue less the weight of filter nah.

Determination of Magnesia.—The combined filtrate from the calcium exalate precipitate is boiled with 1 to 2 g. of sodium ammonium phosphate; about 10 e.e. of concentrated ammonia are added and the solution is allowed to stand in the cold for at least 3 hours. The mixture is then thoroughly stirred and filtered and the precipitate washed once with cold water while the filtrate is rejected. A hole is made in the apex of the filter paper, the precipitate washed into the beaker from which it has just been filtered and the magnesium ammonium phosphate dissolved in a few e.e. of nitric acid, in the way described under the determination of lime. The solution is again precipitated, exactly as indicated above. The cold solution, after standing, is vigorously stirred, filtered through an ignited and weighed Gooch crueible and washed with a cold 10% ammonia solution until a few c.c. of the washings give no precipitate with a nitrie acid solution of silver nutrate. The precipitate is dried, ignited, cooled, and weighed as magnesium pyro-

chlorides which interfere with the determination. | sulphate—the conversion factor to magnesium oxido being 0.3621.

Estimation of Alkali Oxides.—0.5 g. of tho very finely powdered dry substance is thoroughly ground up with about 0.5 g. of ammonium chloride and 3 g. of calcium carbonate in an agate mortar, and the mixture transferred to a platinum cruciblo. The mortar is "rinsed out" with another gram of calcium carbonate and the "washings" added to the mixture in the crucible. After well stirring the contents with a glass rod, the crucible is heated over a small Bunsen flamo for about 15 minutes, during which period the erueible is partly covered by the lid. When all the ammonium salts have volatilised the lid is placed in position and the lower third of the erucible heated to dull redness for I hour, but the cake must not be fused. The crueible is cooled, the cake transferred to a porcelain basin and the crucible thoroughly rinsed out into the basin with hot distilled water. After a few minutes the cake is gently powdered in the basin with an agate pestle; the mass is then digested with about 80 c.c. of hot water for 30 minutes, filtered and tho residue washed with hot water until the washings occupy 120 to 150 e.e. The residue must bo well washed, particularly if the amount of alkalis is large. The residue is rejected.

The filtrate is heated to 80° to 90°C. and 10 e.e. of ammonium carbonate solution I added to precipitate the lime. The liquid is filtered and the filtrate retained. A hole is made in the apex of the filter paper, the precipitate washed through into the beaker from which it was filtered and redissolved in about 5 e.e. of hydrochloric acid. The solution is boiled, a slight excess of ammonia and ammonium carbonato added and the liquid filtered into the vessel containing the first

filtrate, the residue being well washed.

The combined filtrates are evaporated to dryness in a large platinum or silica dish. The dish is then covered with a clock glass and baked for an hour at 120°C, in an air oven. After removing the elock glass, the dish is ignited at a dull red heat until all the ammonium salts have volatilised. The cold residue is moistened with about 3 c.c. of ammonium oxalate solution, in order to precipitate tho last traces of lime, and allowed to stand for 12 hours, after covering with a clock glass.

The mixture is filtered into a small platinum dish and the residue washed with ammonium oxalate solution. The filtrate is evaporated to dryness on a water bath, but, as the liquid may spurt during the early stages of the evaporation, the dish is covered initially with a clock glass and any liquid adhering to the glass is subsequently washed back into the dish. When dry, the residue in the dish is ignited and, when cold, the contents are moistened with a few drops of concentrated hydrochloric acid, again evaporated to dryness, ignited at a low red heat, cooled and weighed. The residue is then washed out with hot water into a small porcelain basin and the platinum dish again ignited and

weighed. The difference in weight gives the

weight of alkali chlorides.

Unless the reagents are definitely known to be free from alkalis, a blank determination of the alkalıs as chlorides must be made on an equivalent weight of ammonium chloride and calcium carbonate. The amount found, if any, is deducted from the weight of mixed chlorides.

If the silicate contains appreciable quantities obtained partly in the form of sulphates at this stage. In auch cases, before the final pre-cipitation of the lime, the aulphates are converted to chlorides by the addition of a little barrum chloride solution, followed by a few drops of animonium carbonate solution to precipitate

the excess of barrum A few drops of perchloric acid (ap gr 120) are added to the liquid in the porcelain basin, at the rate of 6 cc per g of mixed chlorides, and the solution evaporated almost to dryness on a water bath in an atmosphere free from ammonia fumes 10 e e. of water are added and the solution again eveporated to dry oess The residue is at once treated with alcohol wash liquor sand immediately filtered through a weighed Gooch crucible which has been pre viously heated to 120°C for 14 hours. The residue is washed with about 30 ee of the alcohol mixture and the erucible and contents dried for I hour at 120°C and weighed The weight of the precipitate represents potassium perchlorate which is calculated to potassium ehlonde and deducted from the weight of alkalı oxides are then calculated.

potassium The conversion factors are perchlorata to potassium chloride, 0 5381; potassium chloride to potassium oxide, 0 6317, sodium chloride to sodium oxide, 0 5302

(This section on the snalysis of clays has been written by Mr. H V Thompson, MA) Laterature —The reader should consult original

papers in the Transactions of the Ceramic Society (England), the Journal of the American Cersmic Society, Berichta der deutschen Keramischen Gesellschaft, etc Other works are J. W. Mellor, "Comprehensive Treatise of J. M. Menor, Comprehensive Treatise of Inorganic and Theoretical Chemistry," London, 1935, vol. VI. (The Silicatra), II. Walson, "Ceramics," New York, 1927, Collected Writings of H. A. Seger, Easton, Pa , 1902

CLAY-IRON STONE & CHALLERITE CLAYITE.

A name auggested by J W Mellor in 1909 for the non-crystalline variety of kaolimite, H.Al.Si.O. of which china day and most other clays are largely composed The same name had been earlier used by W. J. Taylor, in 1859, for an uncertain altera tion product (of tetrahedrite!) composed of aulphur, arsenie, antimony, lead, and copper.

Alcohol Wash Liquor -97 volumes of absolute alcohol, 3 volumes of water and 0.25 volumes of per-chloric acid (sp.gr. 1.20). The solution should be the state of the state of the solution and the state of the stoppered place bottle containing a few grams of thely powdered potassium perchlorate. The mix ture is well shaken and filtered into a small dry washing bottle immediately before use.

CLEMATINE or BRILLIANT HELIO. TROPE 2R (r. AZINES).

CLEVEITE. A variety of pranimite, conaisting of a uranate of urany L and oxides of lead and the rare earths. It occurs aparingly as emall cubic crystals in pegmatite veins in the south of Norway; and is of interest as being the mineral in which terrestrial helium was first discovered Hillebrand (Amer. J. Sci. 1890, full of aulphur compounds, the alkalis will be 40, 384) found that on decomposing the mineral with sulphuric acid, "nitrogen" was evolved. Ramsay (J C S 1895, 67, 1107) proved that the gas evolved was not nitrogen but helium, Only half the helium is evolved by heet alone. and, moreover, in some cases the disengacement of the gas is accompanied by considerable evolution of heat (Proc. Roy. Soc. 1898, 64. 1401 See BROGGERITE

L. J, S CLEVE'S ACIDS. 1 Naphthylamine 6

and 7 aulphonie acids. CLIFTONITE : CARBON, GRAPHITE Vol p 3154

CLOVE. The dried flower buds of Eugenia aromatica (Lann) Baill (Fam Myrtacen), a native of Molucca, where it was also formerly enitivated, but now grown chiefly in Zanzibar and Pemba The flower buds are white at 6rst, beroming green and finally red, at which stage they are ready for collection and must be gathered immediately, the time of the year at which the harvest occurs being from August to December The buds are air dried and separated from their peduncles, which then form the clove mixed chlorides to give the weight of sodium stalks of commerce and these are used either as chloride. The equivalent amounts of the asource of oil or to adulterate the buds. Cloves vary in length from 100 mm to 175 mm; the largest and plumpest come from Penang and Amboyna, though very few of these reach the European methet, The Zanzibar clove is brownish black in colour, while that from Penang is reddish brown, and the flower head of the Penang clove appears to be much more fully developed than that of the Zanzibir variety. The whole clove consists of a cylindrzcal calyx tube with four sepals at the top to which the flower bud proper is attached Cloves possess an aromatic odour and a atrong spicy, and pungent taste. Fresh cloves should sink in boiled water, a distinction from exhausted cloves, which are frequently used for adultera-tion. They are used to a considerable extent as a apice and as an aromatic carminative, whilst the essential oil is largely used as a mild pre-servative. Corran and Edgar (JSC.I. 1933, 52, T149) have examined the preservative action, and find that mustard and cloves are for auperior to all other spices, and better than the permitted amounts of sulphur dioxide or benzoic acal for preventing the fernientation of glucose

by yeast Microscopic Appearance.-The chief microscopic characteristics are the numerous oil glands just beneath the surface of the call's and the cally teeth, end also in the petals The pollen grains are tetraliedral and measure from 15 to 20 a in diameter. Itosette, but not presmatec crystals of calcium oxalate present, and the buds contain no starch or

selerenchymatous cells. The bast fibres are

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the proportion of clove stalk to bud may be obtained by counting the ratio of the bast fibres to the isodiametric sclerenchymatous cells in a powder. The stalks also contain prismatic erude fibre. erystals of calcium oxalate.

Chemical Composition .- The following table shows the approximate composition of clove

buds and stalks:

pud pulin and				Buds.	Stalks.
Moisture				£∙8°–0∙3	8.7-10.2
Ash				5.3-7.6	6.9-8.0
Volatile oil				14.0-21.0	5·0-6·0
Fixed oil an	đ	resin	n.	5.0-10.0	3.5-4.0
Protein .				5.0-7.0	5.8-6.0
Crude fibre				6.0 - 9.0	13.0-19.0
Tannin .	•			10.0-18.0	about 18

The principal constituent is the essential oil and this is probably determined most satisfactorily by the method of Coeking and Middleton (Quart. J. Pharm. 1935, 8, 435). The powdered buds are mixed with brino, distilled, and the vapours passed through the top of a condenser into a graduated tube, into which a small quantity of turpentine has been previously distilled to make the mixture lighter than water, the condensed water itself heing returned to the distillation flask. An air inlet and outlet is provided by a side tube between the

bottom of the condenser and the graduated tube. Standards,—The "British Pharmacopæia" requires that cloves shall contain not more than 5% of clove stems and 1% of other foreign organic matter. The ash shall be not more than 10%, the acid-insoluble ash not more than 0.75%. The United States standard requires the buds to contain not more than 5% of stalks, not less than 15% of volatile oil, not less than | given below:

broad, measuring about 50µ, and some idea of 112% of quereitannic acid calculated from the oxygen absorbed by the aqueous extract, not more than 7% of ash, not more than 0.5% of acid involuble ash, and not more than 10% of

Adulteration.—The chief adulterants are clove stems and exhausted cloves. Other adulterants, such as pimento and cereal products, have been

detected.

T. McL. CLOVENE v. CARYOPHYLLENE.

CLOVER. The name given to a number of leguminous crops grown for cattle-feeding. The true clovers are all species of Trifolium and include white clover (T. repens), red clover (T. pratense), alsike clover (T. hybridium), crimson clover (T. incarnatum), yellow suckling clover (T. dubium), mammoth red or zig-zag clover (T. medium), hop clover or hop trefoil (T. procumbens). Other "clovers" of agricultural value are yellow clover or yellow trefoil (Medicago lupulina), Bokhara clover or melilot (M. alba), sweet clover (M. dentatus), Japanese clover (Lespedeza striata), Soola or sulla clover (Hedysarum coronarium), various smaller vetches, etc.

The importance of clovers for cattle feeding lies in the high nitrogen and mineral matter contents. In addition, the soil in which they are grown is enriched in nitrogen as a result of the fixation of atmospheric nitrogen by tho bacteria present in the nodules of the roots. In practice clovers intended for direct grazing are frequently grown in conjunction with "seed" grasses, or if required for winter feeding the unmixed clover is hayed. Average analyses of green clover and hay from (a) European, and (b) American sources are

GREEN CLOVER.

1	Water.	Protein.	Fnt.	Soluble carbo- hydrates.	Flbre.	Ash.
T. repens (a) T. pratense (a) T. pratense (b) T. hybridium (a) T. hybridium (b) T. incarnatum (a) T. procumbens (a)	81.5 79.0 70.8 81.8 74.8 81.5 80.0	3.4 4.4 2.8 3.9 2.8 3.5	% 0.8 0.7 1.1 0.7 0.9 0.7 0.8	% 6·9 9·4 13·5 7·0 11·1 7·0 8·4	%3 5.9 8.1 6.2 7.4 6.1 5.7	2:1 1:6 2:1 1:5 2:0 1:9

CLOVER HAY.

	Water.	Protein.	Fat.	Soluble carbo- hydrate.	Flbre.	Ash.
7	9.7	2; 15.7	2.9	39.3	24°1	8.3
T. repens (a)	16.5	13.5	2.9	37.1	24.0	6.0
T. pratense (b)	15.3	12·3 13·6	3·3 3·1	38·1 31·5	24·8 25·7	6·2 7·1
T. hybridium (a) T. hybridium (b)	16∙0 9∙7	12.8	2.9	40.7	25.6	8.3
T. incarnatum (a)	16.7	12-0	2.4	35.5	26.2	7.2
T. procumbens (a) T. rnedium (b)	16·0 21·2	15·4 10·7	3·4 3·9	33·2 36·6	24·5 24·5	7∙5 6·1
, Laspedeza etriata (a)	9.1	13.7	4-0	47.5	21-6	4.1
Lespedeza etriata (b)	11.0	13.8	3.7	39-1	24.0	8-5

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as 40-50% may be in non protein or amide forms in 85-90% of in young plants. In mature plants of the nitrogen is present as protein. In New Zealand and elsewhere the presence

poisoning of cattle or even any ill effect on milk dry matter.

Of the total nitrogen content of clovers as much flavour is likely to be of rare occurrence. Armstrong and Horton (Proc. Roy. Soc. 1913. B. 86, 262) detected an enzyme capable of hydrolysing cyanogenetic glucosides in certain varieties of wild but not of cultivated white

clover. of evanogenetic glucosides in clovers has caused some anxiety. According to the investigations of Doak (New Zealand J. Agric, 1935, 5t, 159) Wolff's analyses of the ash of clovers include the following, expressed as percentages of the

	Total ash	к,0	N210	cio	HgO	Fe ₁ O ₃	P205	202	510,	CI
T. pratense T. incarnatum T. hybridium T. repens	6 83 6 08 4 76 7 16		0 14 0 52 0 15 0 54	1 92 1 62	0 37 0 60	0 07 0 12 0-02 0 17	0 43 0 48	0.15	0 16 0 99 0-19 0 30	0-27 0-22 0-26 0-26

A. G. Po.

CLOVER FLOWERS. It has long been II contains a methoxyl group and forms a known that clover flowers dye a yellow colour tetra-actyl compound, colourless, glatening on aluminum mordanted fahries, and in the primatic needles, in p. 145-147. past they were employed to a minor extent for dyeing purposes. Three varieties have been m.p. 210°, dissolves in alkali carbonstes yielding ehemically examined. Trifolium pratense, T. yellow solutions, and with alcoholic ferms incarnatum, and T. repens

Trifolium pratense —According to Power and Salway (J C S. 1910, 97, 231) the flowers known "common red clover" contain, in as the addition to isorhamnetin :

and a glyconde of querectin, m p 235°, numerous other phenohe substances which are described below. These appear to be closely allied to the

colouring matters of the flavone group.

Pratol, C₁₉H₈O₂(OH)(OCH₈), colourless needles, m p. 253°, readily soluble in hot aqueous sodium carbonate and sodium hydroxide with a pale yellow coloration, yields a mono-acetyl derivative, feathery needles, mp. 166°. Robinson and Venkataraman (J.C.S. 1926. 2344) observe that pratol closely resembles and is very probably identical with 7-bydroxy-4'methoxyflavons.

m p. 262° (acetyl derivative, feathery needles, m p. 167°-168°), prepared by heating resarctophenons with anisie anhydrids and sodium anisate and hydrolysing the product.

A compound, C14H15O7, thin yellow plates, m.p. about 280°, is soluble in alkalis with a yellow colour and gives with sulphure acid a solution exhibiting a hrilliant green fluorescence. following a hrilliant green fluorescence.

Pratensol, C1,H,O2(OH), feathery needles, chloride gives a greenish-hlack coloration. Triacetyprateriol, colourless, slender needles, melts at 189°. Robinson and Shinoda (J.C.S. 1925, 127, 1973) suggest that pratensol may be a

trihydroxystyrylchromone related to apigenin. A phenolic substance, C₃, H₁O₂(OH), color less needles, m p. 225°, is soluble in alkali bydroundes forming colourless solutions, and gives with alcoholo ferrio chloride a dark green coloration. The acctyl derivative, alky peedles, melts at 209°

The glycoside trifolin, C., H., O, p. ls yellow needles, m p. about 260° (decomp), s coluble in alkalis with an intense yellow colorstion, and dissolves in sulphuric acid forming a yellow solution which rapidly develops brilliant green fluorescence. When hydrolysed at yields rhamnose and trifolitin, C18H (O (OH)), slender reliow needles, mp about 275° slender yellow needles, mp (decomp.). Alkahs dissolve trifolitin with an gives a dark green coloration, and alcoholo basic lead access asic lead acetate an orange yellow lead saltthe contains no methody group and is maltered when heated for several hours with 30% aqueous potassium hydroxide. It does not appear to belong to the flavone group, and differs from these by the fact that it does not give an exemum salt with sulphuric acid and only with difficulty a potassium compound by treatment with alcoholic potassium acetate. It may possibly consist of a tetrahydroxyphenylnaphthaquinone. The tetra-acetyl comound, colourless silky needles, when rapidly heated melts at 116°, resolidifies at a higher temperature, and finally melts at 182°.

The glycoside isotrifolin, C2:H2:O11, pale yellow needles, m p. about 250°, is isomeric with trifolm, and when bydroly sed yields, similarly to Though in general the latter, trufolitin.

behaviour it is very similar to trifolin, it is much more soluble in alcohol, and does not appear to be identical with this glycoside.

In addition to these compounds, the flowers contain salicylic acid, coumaric acid, myricyl alcohol, C₃₁H₆₃OH, heptacosanc, C₂₇H₆₆, hentriacontane, C₃₁H₆₄, sitosterol, C₂₇H₄₆O, trifolianol, C₂₁H₃₅O₂(OH)₂, palmitic, oleic, lineleic, and isolinolenic acids.

Trifolium incarnatum.—A considerable difference is exhibited between the constituents of the "carnation or crimson clover flowers" and those of the T. pratense or "common red clover." According to Rogerson (J.C.S. 1910, 97, 1001) these flowers contain pratol, quereetin, and a gluceside of quercetin termed incarnatrin, C₂₁H₂₀O₁₂,3H₂O, yellow prismatic needles, m.p. 242°-245°. Incarnatrin dissolves in sulphuric acid, forming a green fluorescent solution, and when hydrolysed yields querectin and glucose. It is isomeric but not identical with the quereimeritrin of Perkin (J.C.S. 1909, 95, 2181).

In addition to these substances the flowers yield furfuraldeliyde, benzoic, and salicylic acids, a trace of p-countaric acid, incarnatyl alcohol, C₃₄H_{c9}OH, hentriaeontane, a phytosterol, C₂₇H₄₆O, and palmitie, stearie, oleic, liuoleic, and isolinolenic acids.

Trifolium repens .- The flowers of the white clover, T. repens, according to Perkin and Phipps (J.C.S. 1904, 85, 58), owe their tinetorial property to querectin which is present as glucoside.

Nakaoki (J. Pharm. Soc. Japan, 1933, 53, 238) isolated trifoliin from the flowers and considered this to be a quercetin rhamnoside. According to Hattori, Hasegawa, and Hayashi (Acta Phytochim. 1937, 10, 147), however, trifoliin is identical with isoquereitrin (quercetin 3-glucoside). A. G. P.and E. J. C.

CLOVES, ESSENTIAL OIL OF. Distilled from the unexpanded flower buds of Eugenia caryophyllata Thinb. (Fam. Myttacew), a native of the Molucca Islands and cultivated in Zanzibar, Pemba, Amboyna, Penang, Madagascar, and in the Seychelles, Réunion, Ceylon, and Mauritius. The yield of oil from the buds is 16 to 19%. The ripe fruits (mother cloves) and the stems yield 4 to 7% of oil slightly inferior in quality. The leaf oil is also distilled in Mauritins.

Constituents.—The chief constituent is the phenol cugenol, of which 80 to 92% is present. Engenyl acetate occurs to the extent of about Other constituents are furfural, methylfurfural, dimethyl furfural, methyl amyl ketone, methyl heptyl ketone, methyl benzoate, methyl amyl carbinol, methyl heptyl carbinol, methyl and benzyl benzoates, methyl salicylate, and the resquiterpenes a and \$-caryophyllene.

Characters.-A colourless or pale yellow oil, darkening on keeping. Sp.gr. 1.047-1.065, opt. rot. at 20° 0° to -1.5° , $n_{\rm D}^{20}$ 1.528-1.540, phenol content 82 to 92%. Soluble in 2 volumes of 70% alcohol. The phenol content is determined by adsorption with cold 5% aqueous rolution of potassium hydroxide as described in the "British Pharmacopecia."

OXYGEN-CONTAINING COMPOUNDS.—m. and p-content is determined by adsorption with cold 5% aqueous rolution of potassium hydroxide as described in the "British Pharmacopecia."

OXYGEN-CONTAINING COMPOUNDS.—m. and p-content is determined by adsorption with cold 5% aqueous rolution of potassium hydroxide as described in the property of the cold of the property of the

CLUPEINE is the protamine of herring sperm. It contains 15 amino-acid residues and 14 peptide linkages and yields on hydrolysis arginine (10 mols), scrinc (2 mols), prolinc,

alanine and valine (1 mol. each); mol. wt. 20 Z1.

COAL TAR.—The dark brown to black, oily to viscous product of characteristic odour obtained as a by-product of the carbonisation of coal in gasworks retorts or in coke ovens (see also TAR).

Coal tar is a colloidal system, the disperse phase consisting of coarse to ultramicroscopic particles, the so-called "free earbon," as determined by the matter insoluble in aniline or pyridine, the continuous phase being the complex mixture of solid and liquid chemical compounds which form the tar oils. According to Nellensteyn, resin-like hydrocarbons adsorbed by the free carbon act as protective colloids to confer stability on the system. Upwards of 120 compounds have been identified in coal tars formed at high temperatures, i.e. over 900°C. In the fellowing list those shown in CAPITALS are produced commercially, those for which commercial methods of isolation bave been devised but not yet applied on a large scale are shown in black-face type. The compounds are given in the general order of their boilingpoints, but in some eases the compound may be found in a higher fraction than that indicated by its boiling-point owing to the formation of double compounds.

Boiling-point below 200°C.

HYDROCARBONS. - n. Butylene and homologues, n-pentane and homologues, cyclopentadiene, di- and tetra-hydrobenzene, BENZENE TOLUENE, ethylbenzene, p., m., o.XY-LENES, styrene, n- and isopropylbenzene, o-, m-, p-cthyltoluenes, mesitylene, pseudocumene, dicyclopentadiene, hemellithene, hydrindene, INDENE and homologues, tetramethylbenzenes, durene.

OXYGEN-CONTAINING COMPOUNDS .--Acctone. CUMARONE and methylcumarone, PHENOL, o.CRESOL.

NITROGEN COMPOUNDS,—Acetonitrile, PYRI-DINE and its METHYL di- and trimethyl homologues, pyrrol, aniline, benzonitrile.

SULPHUR Compounds.—Ethylmercaptan, CARBON DISULPHIDE, thiophen and mono. and dimethyl homologues and ethyl sulphides.

Boiling-point above 200°C.

HYDROCARNONS.—Penta- and hexamethylbenzene, methylindene, tetra- and hexa-hydronaphthalene, NAPHTHALENE, α- and βmethylnaphthalenes, dimethylnaphthalenes, diphenyl and its methyl and dimethyl derivatives, 4:5-benzoindane, ACENAPHTHENE, FLUOR-ENE and its methyl derivatives, phenanthrene, fluoranthrene, ANTHRACENE and its methyl and dimethyl derivatives, pyrene, naphthofluorenes, triphenylene, chrysene, truxene, perylene, liquid and solid paraffins.

COAL TAR

β naphthole, diphenylene oxide and its methyl | cresols, is decreased, the pitch yield, naphthalene

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derivative. NITEOGEN COMPOUNDS .- Tetramethylpyridine, unoline and isoquinoline and their methyl denvatives, indole and its methyl denvatives a- and B naphthonitriles, acridine and its methyl and dimethyl derivatives, phenonthridine, a- and B naphthylamine, CARBAZOLE and its 2- and 3-methyl derivatives, phenylnaphthyl carbazole. SULPHUR COMPOUNDS .- Thionaphthen tetra-

methylthsophen. diphenylene sulphide, thiophenole. The properties and composition of coal tars are materially influenced by the type of coal carbonised, by the maximum temperature to which the tar vapours are exposed in the retort, and by the time of exposure to this tempera-The primary tar formed in the retort consists mainly of paraffin hydrocarbons, ole fines, naphthenes, and phenols, and the wide variety of conditions governing secondary decomposition accounts for the range of composition of the products of high temperature carbonisation. The coal components are sob jected to different degrees of heat and the secondary reactions overlap In general, medinm (900°carbonizing temperatures. 1,150°C.) and rapid removal of the tar vapours from the retort give a higher yield of tar (12-14 gallons per ton of coal) rich in paraffinoid hydrocarbons and in high boiling phenols. The arematie compounds present are in part highly hydrogenated and methyl-substituted, while the the contains a high proportion of oils and a low proportion of pitchy matter and of material insoluble in pyridine. Such tars are obtained from vertical retorts, particularly if steaming is employed during carbonisation. In horizontal retort practice higher temperatures are em-ployed (1,150°-1,350°C.) and the time available for secondary decomposition is related to the for secondary decomposition is related to the maximum temperature employed during distilla-density of filling the retort charge. The yield tion and period of exposure thereto (see Adam of tar is normally 9-11 gallons per ton of coal, the content of phenols, now mainly phenol and

content, and matter insoluble in pyridine are increased, and the hydrocarbons are almost wholly of the aromatic series and of a lower degree of substitution. Under comparative carbonising conditions, Midlaod and Yorkshire roals yield tar containing more light oil and less naphthalene and anthracene than Durham coals. In general, young coals have the higher oxygen content and give a higher tar yield. In coke oven practice (temperature 1,000°-1,250°C.) the degree of pyrogenic decomposition of the primary vapours varies with the type of oven and with the time eyele, but generally tar yields are lower at Si gallons per ton of coal, the tar being relatively viscous, highly aromatic in character, the naphthalene content being high and the phenols and pyridine insoluble matter low. Whereas in a vertical retort the vapours proceed npwards and escape through cool inner channels in the coal, in a horizontal retort the vapours travel along the hot upper surface and are exposed to radient heat. In a coke oven the large mass of wet coal cools the walls and the tar in part condenses in the centre of the coal mass and later, as carbonisation proeceds, is again vaporised, the degree of further decomposition depending on the temperature of the oven top. Pyridine insoluble matter (elementory analysus-C about 90%, H about 3%, balance O+ N+S+ash) is the final product of decomposition of the tar vapours local heating occurs, the absolute amount of this matter does not increase during the subsequent distillation of the tar. The percentage of matter insoluble in toluene but soluble in pyridine is relatively constant in crude tars, but increases in amount during the tar distillation process, the formation ratio depending on the prior thermal history of the tar, and on the and Sach, J.S.C.I. 1929, 48, 337T).

Typical properties of crude coal tars are:

Borizontal Vertical retorts. Coke ovens. retorts Specific gravity at 15 5°C. 1-12-1-18 1-16-1-23 1 07-1-12 Distillation per cent, by wt. of dry tar: Light oil to 170°C. . Middle oil to 230°C. 1-3 3-6 5-10 7-12 11-15 8-15 Creosote oil to 270°C, 10-15 11-20 Anthracene oil to 350°C. (approx.) 15-24 11-14 10-16 Patch (medium grade) 68-67 45-51 47-65 12-23 3-B 5-15 Matter insoluble in toluene 3-7 Phenols by per cent, volume on dry tar 3-6 6-10

The pitch from a horizontal retort tar will i contain 22-35% matter insoluble in tolucce 16,500-17,000, net 15,900-16,400 B.Th.U. per (free carbon) and will yield 53-70% volatile 1b. The specific heat is 0.35±0.05 at 400 matter, that from vertical retort tar will contain 045±0-05 at 200°C. ("International Critical 10-20% free carbon and yield 67-75% volatile Tahlea"). Spiers (Technical Dats on Firel, 1933 matter. Coke-oven pitch gives on analysis quotes as a typical ultimate analysis, C 860% 15-25% matter insoluble in tolures and 63-72% | H 6-2%, N 18%, S 1-6%, O, ash, and errors 50%.

volatile matter. Wess and Downs (Ind. Eng. Chem. 1923, 15, 1022) give results of a full invratigation of the respectively as follows: 85 6, 11-4; 88-0, 68; constituents of a cole oven tar.

The calorific value of crude tar is gross Tahlea"). Spiera (Technical Dats on Fuel, 1935) H6-2%, N1 8%, S1-0%, O, ash, anderrora 5-0% The carbon bydrogen ratios of low temperature, vertical, coke-oven and horizontal tars are 890.55: 915.52.

specific gravities at 15.5°C, of the fractions are shown in the following table:

Fraction.	Horizontal tar.	Vertical tar.	Coke-oven tar.
230°-270°	1.03-1.01	0.990-1.00	1.035-1.05
270°-300°	1.05-1.065	1.00-1.03	1.06-1.07
300°-350°	1.095-1.11	1.05-1.075	1.095-1.11

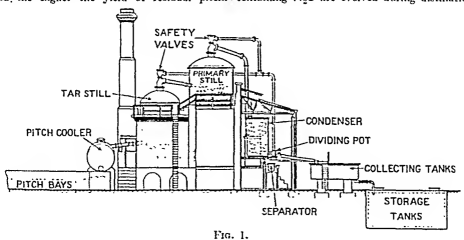
The yield of fine products per ton of crude tar varies within wide limits, and the following figures are to be taken only as a general guide: 90's benzole, $1-2\frac{1}{2}$ gallons; 90's toluble, $\frac{1}{2}-1\frac{1}{4}$ gallons; 90/160 solvent naphtha, 1½-2½ gallons; phenol, 8-15 lb.; cresylic acid, 20-35 lb.; 90/140 pyridine bases, 0.15-0.4 gallon; naphthalene, 20-100 lb.; crude anthracene, 3 to 15 lb.

DISTILLATION.—The crude tar vapours from the retort after condensation are separated from the ammoniacal liquor fraction, and if agitation be avoided during subsequent cooling, the tur as finally separated contains 3-5% liquor by volume. The corrosion of tar stills is attributed to the ammonium chloride present in crude tar acting in conjunction with the resinols, i.e. compounds soluble in aqueous caustic soda, but which can be precipitated from solution in an organic solvent such as benzene by the addition of light petroleum. The heavier corrosive action of vertical retort tars has been found to be due to their higher resinol content (D.S.I.R. Report, 1935/1936).

Chemical changes, generally described as cracking and polymerisation, occur thring distillation, and the greater these changes

When tar is distilled in an Engler flask, the | The extent of the reaction is a function of time and temperature, increasing in some geometric ratio with increase of temperature, and in an approximately arithmetic ratio with increasing time at a constant temperature. Above 345° permanent gases and water are evolved; between 300° and 345° the decomposition is traced mainly by an increase in the pyridine insoluble matter and toluene insoluble matter (Weiss, Chem. and Ind. 1932, 10, 219; Adam l.c). Tars which have been and Sach, subjected to high temperatures during formation are less susceptible to cracking during distillation. These factors, safety and fuel, labour and plant maintenance costs, must be considered in the design of efficient tar distillation equipment.

Crude coal tar, which may be preheated by indirect exhaust steam, is normally distilled in direct-heated wrought iron or mild steel vertical pot stills of 10-40 tons capacity, of height slightly greater than diameter, and having a concavo bottom. The flues cause the combustion gases finally to encircle the walls of the still. Crude tar tends to froth over if rapidly heated, particularly if the insoluble matter content is high; to overcome this trouble, and at the same time to effect an economy in fuel consumption, modern pot still installations are now usually operated on the "double still "system, in which the vapours from the fire-heated still pass through a coil immersed in the erude tar for the next charge. The tar is thus sufficiently preheated to remove the water and naphtha fraction slowly, provided the water content is not higher than 7%. The general arrangement the lower the yield of oils distilling below 350°C. of the plant is shown in Fig. 1. Foul gases and the higher the yield of residual pitch. containing H₂S are evolved during distillation



a furnace and burnt. Cracking during the later stages of distillation is reduced by the admission of dry free steam to the still, and the

and are drawn off through an oxide bed or intermittent pot still depends very largely presed through an explosion prevention box to upon the skill of the man in charge, and the recent tendency has been towards the development of continuous units, which lend themselves to regularity of operation and control of distillato repeated circulation of inert gases such as CO vields, to lower labour costs, to fuel economy by or nitrogen has been proposed to serve a similar heat interchange between vapours and crude purpose, thus increasing oil yields and reducing tar, and to relatively higher output per unit of formation of insoluble matter in the tar (B.P. capital expenditure. For example, a number of 159852). In America horizontal stills have been pot stills, say five, may be connected in series, preferred. The satisfactory operation of the and the tar progressively heated during its 210 COAL TAR.

continuous plant (B.P. 11143, 1911) consista nf a number of cast-iron stills in series, the heating gases passing through W.I. lengitudinal tubes immersed in the tar, This type of plant, frequently combined with a steam-heated Wikner dehydrator (BP. 191295) has been installed in a number of tar distillenes. In the Ah Der-Halden system (B.P. 340370), largely employed in France, the tar, preheated by the outgoing pitch and by the flue gases is fed to a still into which superheated free steam is admitted. The hot gases from the coke faroace pass first over the steam superheating coal, then round the still, and finally over preheating coils. This plant is stated to give a high oil yield, and relatively good fractionation, but the steam consumption at 40-50% by weight on the tar is high. Kahl (B P. 208691, 221773) superimposes a fractionating column on the still and thereby obtains a sharp separation of the frections without re distillation, thus facilitating the production of phenols, accnaphthene, fluorene, carbazole, etc. Owing to the lengthy expesure of the tar in the still to high temperatures, cracking and polymerisation occur and oil yields are low and pitch yields high.
Intermittent and continuous vacuum distilla

tion has also been practised (Raschig, G.P. 250050, Weise, Petcoleum, 1930, 26, 499, 577). The T.I C. process (BP 170617, 184624) takes advantage of the rapid transfer of heat that can be secured by feeding crude tar over the surface of a cushion of molten lead or ticular moment, and production of high slloy contained in a still—the distillation rate is high, giving high oil yields-and the fractional tar to the higher distillation temperatures. condensers give narrow range oil fractions.

This system is used in gasworks with small daily throughput of tar. In the Barrett (U S.A.) system, tar is distilled hy being brought into intimate contact with the hot coke oven gases by mesns of a rotating roller in a continuous still of apecial design (B.P 346753, 347240, 349088, 370387), and the hard pitch produced, which may have a cube in air melting point up to 400°F., is continuously removed and granulated by spraying with water. The hot gases and vapours from the edded to yield narrow range fractions of oil. atill pass through heat exchangers and are then fractionally condensed. Suspended particles of tar in the vapours may be removed by means of an electrostatic precipitator before the vapeurs are passed to the condensing system. A small number of evens on a battery are equipped to deal with the tetal production It was claimed that the process gave maximum eil and minimum hard pitch yield with low operating costs, hence its development in the U.S.A., where, apart from the tar required for road purposes, the main objective of complete distillation is the production of creosete, pitch being a low value by product. The throughput of the still varies with the even cycle time and nes within the limits 100-125 gallens tar dis-tilled per ton of cost earbonised. Cooke and 300°F. Final temperatures are higher than in Holton (B.P. 201645, 416103) use the sensible pot stills, but the time factor is reduced to heat of the hot crude retort gases by circulating below ... the tar in the collecting mains until a dehy drated tar of the required viscosity for road work is two stage pipe still distillation system ilesigned

passage through the plant so that each still obtained; the light vapours pass forward with yields a selected distillate frection. The Hird the gas and are recovered in a static extractor and dynamic extractor in series.

The pioneer user of the pipe or coal still was Lennard (B P. 814 of 1891). His system improved in detail is employed by the South Metcopelitan Gas Co. The preheated cruds tar is primped through a pipe still and at a tempera-ture of 325°C, entera the top of a column into the base of which steam is admitted. Pitch is rontinneusly drawn off from the base of the column and the vapours pass forward through a series of fractional condensers maintained at progressively lower temperatures. In 1907 Wilton (B P. 26910, 1907) erected a cast iron pipe still, in which the tar was heated to 170°C., while being maintained under a pressure of 40 lb per aq in. The hot tar at the coil exit passed through a control valve to a flash chamber at a tmospheric pressure, in which the tar isyer was kept at 4-6 in, depth. The water and light oils flashed off without frothing and the debydrated tar was further treated in pot stills This system was developed to deal with tars of high water and insoluble matter content. It was later adapted for the production of road tars, the debydrated tar from the first coil being passed through a second coil at temperatures up to 275°C and at pressures up to 60 lb. per sq m.
This type of plant is employed by the Gas
Light & Coke Co. at their Beckton works.

The pipe still system of distillation has certain advantages, including relative safety due to the small quantity of tar in the unit at any paryields due to the short time of expesure of the system operates continuously, and sinco 1033 has been further developed by improved deuga of the heat exchangers and fractionsting columns. The rate of flow of the tar in the conditions, thus minimizing the formation of deposits in the pipes and increasing the rate of heat transfer. The heating surfare has been designed to give a uniform rate of heat transfer per unit of surface and finally, hubbling hood towers with controlled reflux ratios have been

Wilton (B P. 307577, 337581, 424645) curculates the preheated tar through a pipe still to a vapour box and part of the residual pitch is passed again to the feed tank, where crude tar is admitted so that the hot pitch effects dehydration of the tar and distillation of the most volatile constituents. The mixture is again circulated through the still, the temperature being raised to 300°C, or higher.

In the American pipe still system the tar a dehydrated in the heat exchanger system and distilled by one passage through the still Plant of this type in U.S.A. vary in capacity from 150 to 450-700 tons per day. Oil yields are 10-15% higher than for batch stills when

Adam end Potter (B.P. 303038) patented

to give a minimum increase of tolurne-insoluble | pipe stills between which is inserted a vaporiser matter and hence high oil yields. Two coils set in one furnace operate under pressure and in series, the pressure being released in a flash box at the exit of each coil. The release of vapours in the second box is assisted by the admission of superheated free steam and the vapours are fractionally condensed in a series of bubble hood columns.

The Koppers Co. have creeted a plant at Ostrava, Czechoslovakia, and another at Beek-

for the removal of light oils and water. tar is heated in the first still to 150°, and in the second to 300°-350°, according to the hardness of pitch required. The volatile products from the pitch column are carried forward and separated into the required fractions in successive bubble hood column stills with the aid of free steam. The first named plant has a capacity of 200 tons and the second 400 tons a day, the fractions taken comprising light oil, carbolic oil, ton, England, designed on similar principles. The | crude naphthalenc, wash oil, and two anthracene tar is distilled in two stages, in two separate oil fractions—see Fig. 2. This system is

FRACTIONATING COLUMNS for NAPHTHALENE OIL ANTHRACENE OIL DITCH CARBOLIC OIL WASH OIL EXPANDER PRIMARY ķ COOLERS CRUDE TAR STORAGE PITCH CARBOLIC OIL NAPHTHALENE OIL

Fig. 2.

units fed to the furnace per ton of tar distilled are 980,000 B.Th.U. in the form of solid fuel, and 96,000 B.Th.U. as steam (from pump exhausts), a reduction of over 25% on average figures for pot stills (Eisler, Zamrzla and Weinkopf, Glückanf, 1936, 72, 184), which are of the order of 130-145 lb. coal per ton of tar distilled.

In another type of plant the hot tar is flashed into a tall bubble hood tower which embodies the evaporator or flash box section and the vapour rises through the trays on which it contacts with descending reflux liquid. At selected points on the tower distillate sidestreams are drawn off.

The composition of the fractions obtained by primary distillation of crude tar varies considerably with the type of plant and type of erude tar. The following data are typical for a modern pipe still (Koppers) and for a double pot still installation distilling a mixed crude tar.

PIPE STILL WITH FEACTIONATING COLUMNS. The crude naphtha distils 95% between 90° and 170°C., sp.gr. 0.885-0.890, yield about 2-3%; the carbolic oil distils 95% between 180°

economical only when a large tonnage is to be tion distils 95% between 211° and 237°, crystalhandled (100 tons per day or over). The heat lising point about 65°C., yield 6-10%; the wash oil distils 95% between 240° and 290°, sp.gr. 1.025-1.040, yield 61%, this fraction also being fluid at ordinary temperatures; the anthracene oil No. 2 distils 95% between 263° and 330°, sp.gr. 1.07-1.08, yield 5%; the anthracene oil No. 1 commences to distil at 323° sp.gr. 1·10-1·12 yield 15-20%.

LIGHT OIL

LIQUOR

Por Still.—Crude naphtha, sp.gr. 0.910-0.960, distils 50-80% within the range 90°-160°, contains 60-65% benzole hydrocarbons, up to 15% naphthalene, and 5-10% phenols, yield 2-4%. The middle or carbolic oil, sp.gr. 0.960-1.005, distils mainly between 165° and 245°, contains up to 40% naphthalene depending on the nature of the tar distilled, and 15-30% phenols, yield 6-14%. The ercosote oil fraction, sp.gr. 1-025-1-045, distils mainly between 220° and 300°C. Under favourable market conditions the phenols may be recovered by a caustic soda wash, otherwise this oil is mixed without chemical treatment with the anthraceno oil fraction to yield timber preserving oils to the British Standard Specifications (see CREOSOTE). The anthracene oil fraction, sp.gr. 1.075-1.11, distils mainly between 270° and 350°, yield 10-20%. In order to preserve the life of the still and and 210', sp.gr. 0.98-1.00, yield 3-6%, content to reduce decomposition, this fraction is obtained of phenole 30-10%, this fraction being fluid at | with the aid of free steam. After condensation ordinary temperatures; the naphthalene frac-lof the vapours, the water is separated and the

COAL TAR.

oil may be cooled to deposit crude anthracene of and 90's tolnole, xylole, and solvent naphtha. about 35% strength. The crude anthracene Until 1930 benzole for use as motor spint was is recrystallised from solvents such as pyridine refined by similar methods. It was then and solvent naphtha to yield a high grade recognised that the sulphuric acid gave rise to

anthracene and erude carbazole.

the pitch residue in the still has the desired softening point when examined by one of the standard methods, the end point being deter mined by the specific gravity or by the appearance of the final runnings or by the temperature of the pitch or vapours. Normally there are three main halts (1) the anft pitch stage (Kraemer-Saruow and stening point, 25°-50°C), the product being employed for various industrial purposes such as hot process tar macadam and pipe coating for prevention of soil corrosion. The pitch at this stage is allowed to cool off to some degree in the still and is then run to containers or to the mixing plant, the soft pitch may be cut back with a selected the order of 0.02% of substances known as creosote oil fraction to yield road tars of the imhibitors. Suitable inhibitors for benzole are required viscosity; (2) the medium and or cresylic acid of high ortho- content or catechol briquetting pitch stage. This pitch has a softening point 70°C ±5° by the Kraemer Sarnow method, or 78°C ±5° hy the Ring and Ball method. This grade is run from the still to coolers which may be storage vessels with an exit to air through a packed tower down which heavy oil is allowed to trickle for condensation of the heavy obnoxious vapours. From the coolers it is pumped while still fluid, either by steam heated reciprocating pumps or hy air pressure, the dry air being admitted through the molten pitch, to overhead cooling trays of 5 tons capacity, or alternatively, it is run by gravity to wrought iron moulds holding 1-21 est, or to pitch beds, where it is allowed to cool into simospheric temperature. The beds are concrete or hrick-lined, and adhesion of the pitch is prevented by hime-warlang the sale of the pitch is prevented by hime-warlang the sale of the pitch is prevented by hime-warlang the sale of the pitch is prevented by hime-warlang the sale of the pitch is prevented by hime-warlang the sale of the pitch is prevented by the pitch is prevented by hime-warlang the sale of the pitch is prevented by the pitch is preven having a Kraemer Sarnow softening point 100°C. and upwards This piteli is east in cooling trays and is used for special industrial purposes, such as clay pigron manufacture, as a reducing agent and neutral oils in the soluble portion and of in metallurgical processes, as a fuel, or it may be further distilled in special stills or in roke ovens to produce rutch coke.

oil distillates are chemically treated to remove phenols and bases. Narrow range fractions are (B.P. 307566, 331542), but have not been then prepared by redistillation for further developed on a large scale for the treatment of

resolution into the pure rempounds.

The fraction to 170° is washed with 10% eaustic soda solution to remove phenols, which are rerovered as such (see Carbolic Acid). and with 25-30% aulphurie acid, crude pyridine bases being recovered from the acid solution hy neutralisation with gaseons ammonia or sodium earbonate. The neutral fraction is then redistilled from a still fitted with closed steam coils, crude benzole, toluole, and solvent a useless by product of the new illuminating su naphtha fractions leing taken These fractions industry. Early uses were as fuel for retorts are refined by treatment with 3-9% by volume in the manufacture of roofing felt, as a paint for of strong sulphuric acid (C.O.V.) to remove wood and stone, and for production of large thiophen, and olefinie hydrocarbons, neutralised blacks. The distillation of tar is first recorded by water and alksh washes, and fractionally in 1822, and in 1830 Anderson at his work in

unnecessary losses of the order of 5-100 The distillation is carried to the point at which due to sulphonation of the aromatic hydroearbons and to the removal of the olefine hydrocarbons which have a definite fuel value. In Germany a restricted acid wash employing 80% sulphume acid was adopted, the treated spirit being distilled through a 60% caustic soda solution to remove sulphonic scidesters. In the country the inhibitor process was adopted in which the chemical treatment is reduced to that necessary for removal of phenols and bases and for reduction of the sulphur content, and to improve the colour keeping properties. The formation of gum from the unsaturated hydrocarbons during storage is prevented by the addition after redistillation of small amounts of (Hoffert and Claxton, J.S C I. 1930, 52, 25T).

The carbolic or middle oil fraction may be separated into two main fractions by redistillation and the lower boiling fraction washed for recovery of phenols, or the whole fraction may be so treated (see CARBOLIC ACID; CRESTLIC ACID) The first and serond fractions obtained on redistillation, or the whole fraction, is cooled in pans or in tanks fitted with agitators and the crude naphthalena which crystallises out is separated by draining or centrifuging, and may be further purified by hot pressing. The crude naphthalene so obtained is washed in the molten rondition with 4% sulphure acid 93-96%

The separation of tar into an oil fraction and a pitch fraction by trentment with light petroleum, b p 40°-60°C., and determination of the phenole resinoids soluble in benzene but insoluble in light petroleum, forms the hasis of a useful method for the examination of crude tars. Industrial The naphtha, light oil, and earbolic or middle applications of this method have been patented by Lessing (B P. 130362) and Morgan and Pratt high temperature tars.

COMMERCIAL EVALUATION.—The production of tar from roal was first mentioned by Becker in 1663 and later covered by a joint patent with Steele (B P. 214 of 1681). Clayton (1729) earbonised roal with rerovery of by products the tar being used in place of Stockholm tur From the beginning of the nineteenth century until 1850 coal tar was in the main regarded as redistilled to yield pure and 90's benzole, pure Edinburgh was producing naphths which was

as a rubber solvent. His works also produced pitch and earbon black. The first important bulk outlet was opened up in 1838 when Bethell patented the employment of eoal tar ercosote oil for the preservative treatment of timber for railway sleepers and mine props-this use arose from the rapid decay of the sleepers on the Stockton-Darlington railway, opened in 1825. The light solvents were used as such and as illuminants. Following Hofmann's identification of benzene in coal tar (1845), Mansfield in 1848 laid the foundations of the industrial process for the recovery of benzene from coal tar. From 1856 onwards the development of the aniline dye industry, employing as raw materials benzene, toluene, naphthalene, and anthracene (alizarine 1868), exerted a strong influence. The employment of pitch for coal briquetting in 1842 extended the uses to which the products of primary distillation were applied. The production of tarred roofing felt increased, later to decrease in this country, but was maintained in U.S.A. and Germany. From about 1910 a new factor of immense importance arose with the development of the use of refined tar in road construction, both for surface dressing and for the manufacture of tarnineadam. Topped tars were first used on roads to eliminate the dust nuisance which followed the introduction of the motor car (1896), but later prepared tars were produced which, acting as a binder for the mineral aggregate, gave a road surface possessing the required qualities of stability, durability, and safe riding surface.

The production of crude tar increased in this country and more rapidly on the Continent with the introduction of the by-product oven for the manufacture of metallurgical coke (1876 onwards). A further important step was the introduction in Germany in 1890 of the gas washing process for the recovery of benzole from the coke oven gases, this materially reducing the export of benzene from England to the German dye manufacturers.

The quantity of crude tar produced by the sources:

sent to Macintosh in Glasgow for employment | gas industry has tended to increase, but tho production from coke ovens undergoes marked changes in accordance with the state of the steel industry. Total production, therefore, bears no relation to the demand for the products of distillation, the values for which undergo relatively violent changes as the demand of the respective using industries equals or falls below the supply. The post war employment of the benzole hydroearbons as motor fuels conferred on these constituents a relatively stable price in an unsaturated market. The market values for naphthalene, phenols, and eresols have not been without interest, but the real economic factor for the tar distiller is the price realised for the bulk products, ercosote, pitch, and road tar. Data regarding the production of these products is given in the appro-priate articles. Thus the value of tar as a byproduct depends upon many factors, including the cost of transport of crude tar to the distillery, and of the products to the consumer. Present average values in the gas industry are 21d. to 3d. per gallon erude tar, equivalent to 2s. to $\bar{3}s$. $3\bar{d}$. per ton of eaal earbonised.

On the continent of Europe tar distillation practice follows the same general lines as British practice. In the United States the outlet for pitch is limited and hence about one-half of the erude tar produced, after removal of the more valuable light oils, is burnt as fuel in the adjoining steel works. the tar distillation works, hard pitch for fuel purposes is produced, in order that the maximum yield of creosote oil may be obtained. As illustrating recent American conditions, tho following figures supplied by S. R. Church are interesting: "The revenue per American gallon of coke oven tar fell steadily from 5.1 cents in 1929 to 3.7 cents in 1933, and had

recovered to 4.1 cents in 1935.

Complete production statistics for Great Britain are not readily available, but the following data for 1936 have been assembled from the Annual Reports of the Chief Inspector of Alkali, etc., Works, England and Wales, and from other

Origin of Tar.	Tar distilled.	Tar distilled. Pitch pro- duced. Pitch pro- exporte		Tar treated for production of road tar.	Road tar exported.
Gas werks	tons. 1,060,000 709,900	tons. 275,200 260,500	tons.	70	tons.
Other works	20,000 1,789,900	7,600 543,300	407,459	43	169,984

173,016 tons; pitch produced,

1.435,000 tons, of which approximately four-

The figures for Scotland are: Tar distilled, in addition to 171,000 tons of road tar imported. 47,707 The annual production of tar in the United States from horizontal and vertical retorts is The production in Germany for 1935 was about 220,000 tons, and from coke ovens, including those on gasworks, was 2,017,500 tons in 1935 and 2,588,000 tons in 1936; in 6fths was from coke ovens; the road tar contons in 1935 and 2,588,000 tons in 1936; in sumption was about 159,000 tons. In 1936 1935, 465,000 tons were used for road purposes. the production rose to 1,680,000 tons. The The quantity of tar distilled in the United French production of erude tar, approximately States has increased in recent years—the figure one-half from coke ovens, was 525,000 tons; for coke oven tar in 1932 being \$11,500 tons and 305,000 tons of tar were used for road purposes in 1935 1,316,000 tons, the balance of production 214 COAL TAR.

in each year being either hurned as fuel in oxide) containing a variable quantity of strel works or sold as fuel.

the order of 7-8 million tans.

ANALYSIS -Standard methods for testing tar and its products are fully described in the book assued by the Standardssation of Tar mispickel, in sulphide of lead, in cerite, and in Producta Tests Committee (2nd ed., 1938), some peats and coals. which should be consulted. Methods for

British Standard Specification 616/1938. The following determinations are normally made on a sample of crude tar : Specific gravity, water (by distillation), matter insoluble in toluene, ash, and distillation test, followed by the estimation of phenols and of crystallisable solids in the fractions. For the distillation test, the flask containing about 250 c.e. tar is heated in a fusible alloy bath and fractions to 210°, 230°, 270°, 300°, and 360°C, are collected, and the phenols, taken as the volumetrie perrentage of the fractions recorded Alternatively, distillation may be conducted in a tared steel still, about 5 litres of the sample being taken, fractions being collected as before. Pyridine bases and phennis are extracted from the oils distilling to 2707, the former by 25% sulphune and also near Port Macquarie in New South acid, and the latter by 10% aqueous caustic Wales Cobalt has also been obtained from the color of mined by neutralising the respective adutions The oil fractions to 270° are cooled to 155°C and the naplithalene which separates is trans ferred to a Buchner funnel, pressed, and weighed. Similarly, the crude anthracene which separates from the oils distilling between 300° and the pitch point is determined. The pitch remaining in the still is weighed and the softening point, which should normally be 65°-75°C, by the Kraemer Sarnow method, is determined. The washed light oils fractions are redustifled and the benzole (to 100°), tolucie (to 120°), solvent naphtha (to 160°), and heavy naphtha (to 190°) fractions are measured (see also PITCH, ROAD

Literature - A. R. Warnes, " Coal Tar Distillation, "3rd ed.; Lunge, "Coal Tar", Rosendabl, "Steinkohlenteer", J. M. Weiss, Chem. and Ind. 1932, 10, 219, 216, E Weiss, Petroleum, 1930, 25, 499, 577, "Standard Methods for Testing Tar and ita Products "1938; P. Spel-mann, "The Constituents of Coal Tar"; Longmans, Green & Co , British Standard Specifi-

cations (various).

COBALT Sym. Co, at, wt. 58 94, at uo. 27,

1sotopes 57, 59 Occurrence.-Cobalt usually occurs combined with arsenic or sulphur, and is almost invariably associated with nickel and other metals. Metallie cobalt occurs in meteorities The most plentiful and important ore of cobalt is smallite nr tin white codall, GGAS, which usually contains some nickel owing to the presence of chloralite, NIAS, and also iron Sulpharsenide of cobalt, CoAsS, occurs as cobaltile nr cobalt glance. As hydrated arenate, Co_As_O_BH_o\circ tha metal as results in the formation of a matter or subhable found in erythrite or cobalt bloom, and as containing the cobalt which also is subsequently aulphide it occurs in linnesie or coold parites, treated for the separation of the cobalt.

Co, S, Asbolite, asbolan, or black earthy During the amelting for apers or n

cobalt oxide mixed with it, sometimes amount The total world production of crude tar is of ing to 40%, sulphide of cobalt and order of coper, nickel and iron are frequently present. Cobalt occurs in small quantities in many other minerals-in pyrites and rertain iron ores, in

The most important source of cobalt is sampling are given in this publication and in Northern Rhodesia, where it occurs as colabiferous copper ore in the N'Kana mine of the Rhokana Corporation; next in importance comes the Belgian Congo, where it also occurs as cohaltifecous copper are in the Katanga region, followed by the Cobalt district in Ontario, where smaltita and cobaltite are found abundantly in the silver ores, the cobalt minerals running in veins. In Sudbury, Ontario, cobalt occurs as cobaltiferous pyrrhotite associated with nickel, and is recovered as a by product, Other sources are Burma, where nickel spens carrying 3-40 cobalt is a by product from the Burma Corporation amelting operations, French Moroeco, where it occurs mainly as smaltite, and China, where it occurs as cobaltite and amaltite. Asbolite occurs in New Calcionia Central India, Tunaberg in Sweden, Narimanov in Russia, and Cornwall,

EXTRACTION -The methods adopted for the extraction of cobalt from the ores vary with the nature of the ores, and as these generally contain larger quantities of some other metal than cohalt this latter is obtained as a hy-product in the form of a spease containing also arsenide of other metals, as a matte containing sulphides of other metals, or as a precipitate during wet treatment of ores.

Experiments on flotation concentration have given some interesting results. Starting with an ore containing 15 81% Co associated with pyrites and sphalerite, the Canadian Government Laboratories have succeeded in getting a 97% recovery of a 24 77% conventrate, and an 85% recovery of a 29 5% concentrate by flotation (J. S. Godard, Canada Dept. Mines, Mines Branch, Rep 728, 1933, 75-76).

When cobalt ores are treated direct, they are smelted in low-pressure blast furnaces or cupoles together with fluxes which vary according to the nature of the minerals present. When F. M. P. silica is present in excess, lime is added, and when lime or alumina is in excess ailica or a siliceous ore is added to the charge. If the alag obtained be too viscous, then fluorspar is also added. When arsenides are thus smelted a spess or arsenide is formed, which is tapped from the furnare at certain intervals for subsequent treatment. The slag obtained may con tain I to 2% cobalt, it is re-treated in the furnare by which means its cobalt content may be reduced to 0.5% or less. With sulphide ores, which are uncommon, a similar treatment

During the amelting for spess or matte, 1 cobalt is essentially word (hydrated manganese certain amount of iron is allowed to go into the

of cebalt able to pass into the slag, and the iron is also useful in the subsequent separation of arsenie, and in the manufacture of ferrocobalt.

The speiss as obtained is crushed, finely ground, and roasted in a mechanically rabbled furnace, the arsenious oxide which is driven off being condensed in long flues or in some form of dust precipitation chamber, collected and resublimed for sale as white arsenic of commerce. This reasting is continued until the reasted mass contains 10% arsenic, when it is removed from the furnace and again ground. The material is treated with hydrochloric acid and licated and agitated by means of live steam; the cobalt, nickel and copper exides present are converted into chlorides, and the chlorino evolved oxidises the ferrie arsenide to ferrie arsenate, which, being solublo in hydrochloric acid, remains in solution. The solution is allowed to settle and is then filtered, any insoluble compounds including siliceous matter being left behind as a residue. The solution is nearly nontralised with milk of lime and the ferric arsenate is precipitated. Should there bo insufficient iron present to convert all the arsenic into ferric arsenate, iron is added to the solution as ferric chloride. Any arsenic not precipitated by this method, together with any copper present, is precipitated by gassing with hydrogen sulphido; if the copper is high in quantity, it is separated electrolytically. Any iron not previously precipitated is removed by treating the solution with milk of lime until noutral to litmus and adding bleaching powder solution, then filtering off the ferrie hydroxide. The solution thus obtained contains cobalt and nickel chlorides, together with traces of impurities, and is of a fine port-wine colour. It is treated with milk of lime and bleaching powder solution which precipitates the cobalt as hydrated oxide, $Co_2(OH)_6$. With care, this operation can be so adjusted as to stop at a point where the red colour of the solution just disappears, leaving a green colour characteristic of nickel chloride, thus:

$$\begin{array}{ll} 2 \text{CoCl}_2 + \text{CaOCl}_2 + 2 \text{Ca}(\text{OH})_2 + \text{H}_2\text{O} \\ &= \text{Co}_2(\text{OH})_4 + 3 \text{CaCl}_2 \end{array}$$

The green-coloured solution is treated with milk of lime, when the whole of the nickel is precipitated as green nickel hydroxide (T. H. Gant, Chem. and Ind. 1925, 44, 157).

The black precipitate of cobalt hydroxide obtained as above is transferred to tubs, where it is boiled with dilute hydrochloric acid for the removal of traces of nickel. It is then filterpressed and roasted, being finally wet-ground, again treated with dilute hydrochloric acid, washed to remove soluble lime salts and further traces of nickel, filter-pressed, and again roasted. when it is dehydrated and converted into the oxide, Co2O3, which is ready for the market as such or for reduction to the lower oxide or to metal.

When the product of the first smelting is a matte, as from sulphide ores (New Caledonia), this matte is carefully roasted at a moderate of the oxido and charcoal under a layer of temperature only and the sulphides of the metals suitable glass; the fusion is preferably

product because its presence reduces the amount | present, iron, copper and cobalt, thus oxidised to sulphates. These sulphates are dissolved in water and the iron present is precipitated by the addition of the requisite quantity of sodium carbonate. The copper and any licavy metals present are separated by hydrogen sulphide, which is filtered off. After boiling off the excess of hydrogen sulphide the cobalt is precipitated by the addition of sodium hypochlorite (bleaching-powder cannot be used owing to the presence of sulphates in the solution), and the nickel in the remaining solution is precipitated by sodium earbonate.

At La Panda (Belgian Congo) the cobaltiferous copper ores are first smelted for copper in a blast furnace and the slag, which contains about 6.5% Co, 5% Cu, and 10% Fe, is charged with a calcarcous flux and a little coke into open single-phaso electric furnaces taking 6,000-8,000 amp. The product is an alloy containing approximately 28% Co, 30% Cu, and 40% Fe, together with a slag which is returned to the copper smelting furnace. The upper electrode is of graphite and the lower electrode is embedded in the conducting hearth of the furnace. The furnace takes 50-eyele A.C. at 60 volts and consumes about 2,000 kw.-hr. and 8-9 kg. of electrodes per ton of alloy produced (R. Sevin, J. four élec. 1927, 38, 17, 211). This alloy is exported to Belgium, where it is treated for the recovery of cobalt and copper. It is crushed and treated with sulphuric acid so that tho cobalt and iron pass into solution and the copper remains as a metallic spongo. Iron is removed from the solution by the addition of lime and blowing in air, and after further purification the resulting cobalt sulphate solution is electrolysed (J. Lecmans, Chim. et Ind. 1927, 17,

Until recent years the only market for cobalt compounds was in the ceranic industry, but tho metal itself is now largely used for electroplating and as a constituent of many alloys. The cobalt sesquiexide, Co2O3, produced as above, when heated to a high temperature, yields the monoxide CoO. Metallic cobalt is produced by heating the oxide with carbon or carbonaceous material, starchy matter often being used; in this latter case the mixture is made into a paste, shaped, and cut into cubes which are heated to a high temperature in a retort. The metal so obtained is not melted down and contains a certain amount of free carbon. Cobalt is also produced as a grey metallic powder by heating the oxide in a current of hydrogen; if too low a temperature be used (250°) the resulting metal will be pyrophoric, igniting and becoming reexidised in centaet with air, the ignition temperature being 3° only. Carbon monoxide readily reduces cobalt oxide at temperatures above 500°, the reduction being very rapid at 900°. Reduction with aluminium by the thermit process gives a metal containing about 0.1% aluminium (Kalmus, J. Ind. Eng. Chem. 1914, 6, 107). Pure cobalt may be obtained as a powder by heating cobalt oxalate, Co(CO₂). A button of the pure metal may be obtained by fusing the oxalate alone or a mixture

performed in a lime or magnesia crucible at the Co. (CO), are formed which melt at 51°, are highest temperature of the furnace. Pure soluble in alcohol or ether but insoluble in cobalt may also be obtained by electro deposition water, and decompose above 52° yielding from a suitable electrolyte and by heating cobalt Co(CO), which further decomposes at 10° carbonsl

Properties-Cobalt is a brilliant silverwhite metal with a slight bluish tinge. It ranks next to trou as a magnetic metal and retains sulphate of cobalt and ammonium. Deposited its magnetism at high temperatures; above 1,100°-1,150°, however, it becomes para-magnetic. There are two alloteopic modifications of cobalt a cobalt exists at ordinary temperatures and has a stable, close-packed, hexagonal lattice, but at 400° \$ cobalt is formed, which is stable above this temperature and possesses a face-eentred cubic lattice. Both varieties may exist side by side at ordinary temperatures, but a cobalt is the stable form and 8 cobalt the unstable form. Cobalt melts at 1,480° and boils at 2,375°/30 mm. Its specific gravity is 8 756 as east, 8 81 after annealing, and 8 925 after awaging. It is harder and more teracious than iron, its Brinell haedness number varying between 124 and 130, its tensile strength being about 34,400 lb per aq in in the east state, 36.9 0 lb in the annealed condition, and 100,000 lb after colling, and its compressive strength as east 122,000 lb per aq in, after annealing 117,200 lb, per rq in Electrodeposited cohalt is harder than the ordinary metal, its Brinell mumber being 270-311, according to conditions for pure copper to 180 for the alloy con (McXaughton and Hothersall, J Electroder taming 90°, cobalt, Cobalt has a mustle Tech Soc. 1930. 5, 63) The electrical results effect on the electrical respectives of copper, ance is 64×10⁻⁴ ohm per cm³ for pure the specific resistance increases rapidly up to metal, and 8 90×10⁻⁴ for 99 73°, samples; 3 2°, coledit, at which figure the resistance the specific heat between 15° and 100° is reaches 945 microbins per cm. at 20°, carbon in the metal increases the tensile and with a minimum temperature coefficient of compressive at rength and the electrical resistance but has little effect on the hardness Pure cobalt may be machined in a lathe as readily as difference of temperature. Copper alloys conpure iron or nickel, but it is somewhat brittle; taining 1-60 cobalt are remarkably ductile, cobalt containing a small quantity of carbon mallcable and tenneious. machines like mild ateel, and whilst pure east cobalt cannot be swaged without cracking, the metal containing earbon may be rolled or awaged down from the cast state provided these operations are earned out on the hot metal It is soluble in hydrochloric and sulphume acids with evolution of hydrogen, and rapidly soluble ability falling off with increasing percentages of in intre acid; it is not attacked by hot or gold, rapidly at first, then more slowly. cold alkali, but if used as an anode in the Zine-With zine, colalt forms a series of electrolysis of alkaline electrolytes it passes solid solutions containing 0.5 to 185% robalt into solution as colloidsl cobaltons oxide. The saturated solution contains 13 10 robalt Cohalt occludes from 60 to 150 times it sow and a cut-cite containing 50% cohalt solding volume of hydrogen, the amount varying at 413°. The compound CoZn, is formed with the mechanical condition of the metal, at 18-9% cohalt with a melting-point of When heated, the powdered metal combines 873°. with chlorine, bromine and iodine. It decomposes steam at a red heat and if finely divided series of alloys of which the freezing-point curre ignites when heated in oxides of mitrogen, and in a current of among so it which the feether and in a current of ammons gas at 470 forms a mirdle. Copy, which decomposes at 600°.

Compact cobalt does not conduct in are at specific continually temperatures, but if becomes approximately the compound of \$50° and \$40° with 250° with 250° and \$40° with 250° with ficially oxidised at a red heat; aulphur unites cobalt. There are two series of solid solutions with red hot cobalt to form aulphide. When containing respectively 68 5 to 90-5 and 90-5 to

(r. CARBONYLS). Cobalt may be deposited as a fine lusteous coating on metals by the electrolysis of a neutral solution of the double on distorate brick, cobalt is a good estalyst for the hydrogenation of etbylene, causing rapid reaction at -20° (Peace and Stewart, J. Amer

Chem. Soc 1927, 49, 2783). DETECTION AND ESTIMATION .- See CHEMICAL ANALYSIS (Vol. II, pp. 553, 578, 580, 612, 656).

ALLOYS OF COBALT.

Copper -- Cobalt forms with copper a series of alloys containing to definite chemical compounds. Two series of solid solutions are formed containing respectively 0-10% and 93.5-100% copper, and alloys between these hunts consist of conglomerates of the two sets of solid solutions, solidification being complete at 1,105°, in these alloys also the magnetic change is lowered to 1,050°; alloys containing 90% copper are magnetic (R. Sahmen, Z anorg Chem. 1908, 57, 1). The Brinell hardness number of the series increases with additions of cobalt to copper from 28 resistance of about 0-00077 and a maximum thermoelectric effect of 33 microvolta for 1°

Gold -Nolten gold and cobalt are miscible in all proportions, the freezing point curve of the senes having two branches mreting at a entertie at 997" containing 90% gold. Solid solutions are formed containing 0-5 5% and 96 5-100% gold All the alloys are magnetic, the magnetis-

Aluminium .- Cobalt and aluminium form a finely-divided cobalt is heated to 150-200° 100% cobalt. There is a minimum on the in carbon monoxide at 100 atm pressure, curve at 1,375° and 0.55% cobalt. The concange-red crystals of cobalt carbon, prompt, pound CoAl on cooling resets with the head

cobalt at 1,165° to form the compound Co,Als. The solid solubility of cobalt in aluminium is less than 0.02%.

Tin.-Cobalt and tin are miscible in all proportions in the molten state and the freezingpoint curve has been determined by K. Lewkonja (Z. anorg. Chem. 1908, 59, 294). This curve shows that when solid, the maximum amount of cobalt soluble in tin is 3.5%. There is a first cutectic at 1,08° and 34% tin and the cutectic line extends from 3.5% to 48% tin, at which point Co₂Sn forms and freezes at 1,151°. There is a second compound CoSn, which forms at 950°; this compound undergoes a polymorphic change at 536°. A second cutectie compound CoSn, containing 66% tin.

Chromium .- Cobalt nnd chromium are mutually soluble both in the liquid and in the solid state. At a temperature of 1,320° the liquidus curve shows a minimum at approximately 47% cobalt. Alloys containing 45-85% chromium undergo a reaction at 1,225°, at this temperature the homogeneous solid solution breaks down into two solid solutions. containing 0-45% chromium show a polygonal structure containing cobalt-rich cores, the chromium increasing from the centre to the outside. In alloys containing more than 55% chromium the chromium content of the grains decreases from the centre to the outside. At room temperatures the alloys containing up to 25% chromium are magnetisable and the temperature at which loss of magnetism occurs decreases rapidly with increasing chromium content, the addition of 10% lowers the transformation temperature from 1,056° for pure cobalt to 686°, 15% to 300°, whilst the addition of 25% lowers it to below room temperature.

Ternary Alloys.—Chromium alloys, together with the ternary alloys containing cobalt, chromium, and tungsten, or molybdenum or the quaternary alloys containing all four of these metals, constitute an important class which are known under the name Stellite, from the Latin stella, for a star, chosen because of the brilliant polish such alloys take, and retain when exposed to ordinary atmospheric conditions. straight binary alloys of cobalt and chromium can be east into bars or other forms and are almost file-liard. They are slightly malleable when cold and distinctly so when hot, and can be further hardened by heating to redness and quenching in water. An alloy containing 70% cobalt, 25% chromium, and 5% tungsten is a distinctly harder alloy which forges at a red heat and takes a good cutting edge; this alloy is suitable for cold working tools such as chisels and wood-working tools. Much harder alloys are obtained by increasing the tungsten to 20%. at which point they can only be forged to a limited extent; and by increasing the tungsten

mass in alloys containing less than 68.5% cast into bars which are subsequently ground and used as cutting tools for steel and cast and again at 940° to form Co₃Al₁₃. A cutectic iron. Molybdenum produces much the same occurs at 654° containing about 1% cobalt. effect as tungsten and the addition of 10% makes an excellent lathe tool. An alloy containing cobalt 45%, chromium 15%, and molybdenum 40% is very hard, cuts glass, and will even scratch quartz. A common range for the composition of stellite is cobalt 45-50%, chromium 25-30%, tungsten 15-20%, and carbon 2.5-2.75%. For cutting tools this may be used as east bars or as tips which are brazed to shanks and blades fitted into slots in milling and boring heads. It has also been used for giving hard tips to picks, for the hard-facing of dies, and for facing aeroplane tail skids and the steel rings of eement kilns. Stelliting is the operation of depositing a layer of the alloy on freezes at 230° with less than 1% cobalt and steel or cast-iron by means of the oxyacctylene occurs in all alloys between pure tin and the torch, the atomic hydrogen torch, or the electric are, to form a hard wear-resisting surface. A new electro-chemical method of cladding ferrous products with stellite is described by R. R. Rogers (Ind. Eng. Chem. 1935, 27, 783). Widia is another cobalt alloy consisting of tungsten carbide containing about 6% cobalt. It is used for heavy-duty lathe work and the turning of glass and bakelite, and is probably the hardest known alloy.

Iron.—Cobalt is an important constituent in

certain steels, especially magnet steels, and when the metal is destined for use in the steel industry it is generally obtained as a mixture of iron and cobalt known as ferro-cobalt. This is marketed in the form of cubes containing from

10-15% iron and 80-90% cobalt.

Nickel.—With nickel it forms a series of solutions both in the liquid and solid condition.

Carbon.—Cobalt dissolves 3.9% carbon at 1,700°, and at the cutectic temperature 1,300° cobalt retains 0.82% carbon in solid solution. The cutectic is found to contain only metallic

cobalt and graphite but no earbide. ELECTROPLATING.—Cobalt is used for electroplating purposes owing to the pleasing colour and resistance to tarnish of the deposited metal. It has been considered that cobalt deposits are harder than nickel deposits, but D. J. Macnaughtan (J. Iron and Steel Inst. 1924, 109, 410) has shown that under varying conditions the Brinell hardness of cobalt deposits may vary between 270 and 311, that of nickel deposits may vary between 180 and 420. A suitable bath contains 565 g. cobalt sulphate (crystal), 20 g. sodium chloride, and 30 g. boric acid per litre. A current density of 100-160 amps. per sq. ft. may be used and the plating may be carried out more rapidly than with nickel (Brass World, 1926, 22, 303).

COBALT AS A CATALYST .- An account of the reactions of olefins in the presence of metals has been given by G. Egloff and C. I. Parrish (Chem. and Ind. 1937, 56, 321) in which cobalt is included. A cobalt catalyst prepared by caleining the nitrate on pumice followed by hydrogen reduction was heated in a current of cthylene (Walker, J. Phys. Chem. 1927, 31, to 25%, giving an alloy containing cobalt 60% 961), no liquids were found up to 545° while chromium 15%, and tungsten 25%, a very hard much carbon was deposited. At 400° there was alloy is obtained which cannot be forged but is

and a small amount of chane, while a very war of the small amount of chane, while a lumina, a nno nuc returned was 97-49, decomposition into a gas containing pink mass; with alumina, a nno nuc returned was 97-49, decomposition into a gas containing the small amount of chane, blue, and with zinc oxide a green (Rinmany). and 130% methane. Reduced cobalt has no effect npon etbylene at ordinary temperature, but above 360° the gas was decomposed into hydrogen, ethane, and carbon with small quantities of benzene and its bomologues. It is less active than nickel but acts similarly (Sahatier and Senderens, Compt. rend. 1897, 124, 616, 1358; 1900, 121, 267; Ann. Chim. Phys. 1905, [viii], 4, 436).

COMPOUNDS OF COBALT. Oxides .- There are three well-defined oxidesof

cobalt, namely, cobaltous oxide, CoO, cobaltosic oxide, Co,O, and cobaltic oxide, Co,O,. Coballous oxide is obtained when cobalt is burnt in oxygen or when oxygen is blown through the molten metal. It is also obtained by roasting the powdered metal in sir, by heating cobaltons hydrexide below 350° in a current of bydrogen, or by heating the bydroxide or carbonate in air. It varies in colour from greenish grey, red, brewn, to black, according crucible. When the desired blue relour his to method of preparation and size of grain developed the mass is ground in water and dired Cobaltous hydroxide, Co(OH)2, is precipitated when potassium hydroxide is added to a solution of a cohaltous salt out of contact with air as a bine or rose-red precipitate, the blue is considered to be CoO,H2O, and the rose CO(OH), the latter gradually turns beated as above.

hrewn in contact with air forming cobaltie According to Hedvall (Arkiv, Kemi, Min, hydroxide Co(OH)3, Cobaltoese oxide is formed superficially when metallic cobalt is beated in air or by beating cobaltic exide or hydroxide in air; when the product is beated in hydrogen it is reduced first to cobaltons oxide of about 250°, and then extracted with water, On heating to 265° it decomposes, leaving Co.O., and at a high temperature yields Co.O. Impure forms of hydrated cohaltse oxide are found in various minerals and the exidation of yields these compounds, the composition of the hydrate depending on the concentration and temperature of the solution and the nature of the oxidising agent used. The preparation of a required temp number of definite bydrates has been reported, 1911, 85, 497). including 3Co.O.,2H,O; Co.O.,2H,O. 2Co.O.,3H,O; 3Co.O.,6H,O; Co.O.,2H,O; Co.O.,3H,O.

and porcelan manufacturers for the production of the finest blue glaze and colour on porcelam glass and other vitrifiable substances. The presence of yzdgs of oxide imparts a bluish clour; the presence of colour in the colour of the colour

green). Zaffre is obtained by beating a mixture of reasted cobalt ore with sand to a temperature below the melting point. (v. infra, p. 219c).
Pigments containing Cobalt, Thenard's

Blue, Cobalt Blue, Cobalt Ultremarine, King's Blue or Azure Blue.—The composition of this pigment varies considerably according to the method of manufacture. The original Thenaed's blue was prepared with phosphete or assenate of cobalt and alumina, but now cohaltous exide and alumina are practically always used. Using the original method, it may be prepared by precipitating a solution of a cobalt sait, neually the nitrate free from iron and nickel, with potassium or sodium phosphate or with aodium arsenate. The gelatinous violet precipitate is thoroughly washed and well mixed with 3-5 volumes of freshly precipitated, well washed alumina precipitated from a solution of alum free from iron by the addition of sodium carbonate. The mixture is dried and calcined at a cherry-red heat in a well revered clay

A similar product may be obtained by the aumultaneous precipitation of the oxides of cobalt and aluminium by the addition of sodium earbonate to a mixture of cobalt nitrate and alnm; the mixed precipitate is washed and

Ocol. 1914, 5, No. 16, 1), cobaltous oxide and alumina are preferably ignited in presence of potassium chloride as a flux. When the rebaltous oxide is in excess the resulting product can be obtained pure by treatment with water to remove the flux and with hydrochlone and and then to metal. Cobolic carde is formed to remove the flux and with hydrochlone and when a concentrated solution of cobolicous to remove the excess of cobal to nice. At 1,100 the bloom to the second of the cobolic carde with potassum chlorate use the bloom alminiate, CoCAI,O, is formed; evaporated to dryness, heated to a temperature at temperatures above 1,100 a green aluminate, 4CoO,3Al,O,, is produced. Wagner and Vanino found that cobalt sulphate could be used instead of phosphate and that the alumina was most suitebly introduced as ammonium alum. To 22 parts alum were added 25 to cobaltous salts in acid or alkaline solutions 5 parts cobalt sulphate and 0 3 parts magnesium or zincsulphate; the mixture was melted in its water of crystallisation and constantly stirred until dry, after which it was heated to the required temperature (L. Vanino, Chem Zig

The Thenard's blue colours include the range of blues known as "Matt" blues. These are prepared by heating cobalt oxida with Cobaltous oxide is used for the preparation of ammonaum alum or potaba alum and alumina the salts of cobalt and also of smalt the glass with the addition of varying quantities of uncotained by heating the oxide with critical order oxide. From 20-30% of uncotable helps to and proreful manufactures to the analyzed order of the colour to the body in the crame and proreful manufactures to the colour to the body in the crame

oxides has an injurious effect on the colour the disadvantage of appearing violet by graphed by this substance; it is therefore light. It is one of the most permanent blue necessary, for the more delicate work, to ensure pigments, being unaltered by acids or alkala-its complete freedom from such impurities.

and is largely used as an oil and water colour,

poisonous and has the advantage of miseibility with other pigments without alteration.

Caruleum, exline or bleu céleste, is a corresponding colour containing oxides of tin and cobalt and calcium sulpliate or silica. It is prepared by heating together cobalt sulphate. tin oxide, and precipitated silica or chalk and its general composition may be oxide of tin 49 66, oxide of cobalt 18 68, calcium sulphate or silica 31-68. It may also be prepared by pre-cipitating a solution of cobalt nitrate with sodium stannate and heating the precipitate as above. It is a fine light blue pigment of slightly greenish tinge, of greater density than Thenard's blue, and retains its colour in artificial light.

New Blue is a blue pigment of shade varying from a pale greenish-blue to a deep turquoise blue, largely used for enamels, it consists of aluminates of cobalt and chromium produced by the action of alum on carbonates and hydroxides of cobalt and chromium, and the subsequent firing of the resulting precipitate.

Cobalt Green, Rinmann's Green .- The constitution of this pigment corresponds with that of Thenard's blue with the substitution of rine oxide for alumina. It has been shown to be an isomorphous mixture of cobalt zineate (CoO.ZnO) with zine oxide which cannot be chemically separated (J. A. Hedvall and T. Nilsson, Z. anorg. Chem. 1932, 205, 425) It may be prepared by mixing a solution of pure cobalt sulphato (or nitrate) into a paste with zine oxide and heating in a muffle furnace for 3 or 4 hours until the desired colour is obtained. Pigments of a more uniform nature are produced by the precipitation of a mixture of solutions of zine and cobalt salts by means of sodium carbonate, phosphate, or arsenate, and heating the precipitate. The phosphate is said to give a purer and brighter green colour than the carbonate.

Rinmann's green is a fine permanent pigment; its colour varies through many shades, according to the proportion of zine oxide present, the darker greens containing less of that substance. It is non-poisonous and unacted upon by dilute acids or alkalis, but is attacked by ammonia It has also the advantage of not affecting, or being affected by, other pigments.

Turquoise Green is another permanent green pigment prepared by firing a mixture of cobalt oxide, ulumina, and chronium oxide, or sometimes the following proportion of freshly preupitated substances are fired: CoCO, 20, Al₂O₃ 40, Cr₂O₃ 20.

Cobalt Bronze is an aminonium cobalt phosphate of a violet colour, with a bronze-like metallie lustre.

Cobalt Yellow, Tischer's salt, aureolm, Indian fellow, consists of potassium cobaltimitrito prepared by the addition of potassium nitrite to a solution of a cobalt salt acidified with acetic acid. Its composition varies with the proportions used and the strength of the solution. Erdmann and Sadtler found that when precipitated from a solution containing much acetic acid, it consists of K3Co(NO2)8

but works better as water colour. It is non-limpurities, unacted upon by cold water or cold acids, and but slowly blackened by sulphuretted hydrogen. It forms a fine pigment for artistic purposes and when used for painting porcelain it produces on baking the usual blue colour, which is of great beauty.

Cobalt Brown is produced by calcining a mixture of aumonium, ferrous, and cobalt

sulphates.

Cobalt Red is produced by heating a mixture of magnesia and cobalt oxide. Solid solutions are formed the colour of which varies with the proportions of the oxides from hight to dark red (Hcdvall, Z. anorg. Chem. 1914, 86, 296).

Smalt, Bleu d'Azur, Bleu de Saxe,-This important substance consists essentially of a potassium silicate glass coloured with cobalt. In the preparation of smalt in Saxony, smaltinto is principally used. In Norway and Sweden the principal ore is cobaltite. The general process is the same for both ores. The selected ore is powdered, freed from the lighter earthy impurities by a process of washing or concentration and roasted in a reverberatory or muffle furnace. The arsenie and sulphur are expelled as oxides, the former being condensed in chambers. The roasting should be continued until only sufficient arsenie is left to combine with the less oxidisable metals, especially copper and nickel, while the greater part of the cobalt is converted into oxide. The product is known as zaffre or zaffler, which is sometimes considered as a criide oxide of cobalt and sometimes as a basic arsenate and is used for painting on poreelam. Towards the end of the process of roasting a small portion is occasionally tested and the process stopped when the finest tinted glass is produced. The material is then ready for fusion for "smalt."

The fluxes used consist of powdered quartz and potassium carbonate. They must be very pure, as the presence of lime or many other substances impairs the beauty of the colour. The proportions of the ingredients vary according to the richness of the ore and the colour required. The potassium carbonate used generally amounts to about one-third of the weight of the zaffre and quartz together. White arsenic is sometimes added to reduce any ferric salt and thus prevent the injurious effect of that substance on the smalt and otherwise to heighten the colour of the product. The exact amount of each substance is found by a test fusion and comparison with a smalt of the required colour.

The mixture is melted down in crueibles or pots made of very refractory clay, free from lime; these pots are strongly heated and the mass fuses in about 8 hours, it is frequently stirred to render it homogeneous and to break the crust which forms on the surface. The quartz and potash react with the formation of potassium silicate, which the cobalt oxide forming the blue smalt, while the mixed arsenides of nickel, copper, and iron with the small quantity of cobalt arsenide which is always present fuse and form a brittle, metallic-looking speies beneath the smalt. with a variable amount of water. It is a bright This usually contains about 3% cobalt and a Millow crystalline powder, very free from considerable quantity of nickel and these are ture for some time without stirring, to allow violet-blue colour characteristic of robalt the speiss to settle, and the smalt is then ladled silicate, or Thenard's blue, a rich turquoise out into vessels of cold water. It is thus blue characteristic of cobalt aluminate, may be cendered granular, brittle, and easy to pulverise. The blue glass is next ground under water and the product passed through a series of depositing vats for sizing, the coarser portions being reground and the finest portions being cemelted if necessary. The marketable portions are dired and sifted. The presence of oxides other than those of cobalt and potash, even in small quantities, exerts a powerful influence on the colour of the smalt. Baryta produces an indugo tinge; sodium, calcium and insepessium produce a reddish shade, iron, a blackish-green very prejudicial to the brighter coloured smalts, manganese, violet; nickel, violet, but less intense; 0 3% are also used in the preparation of ground copper, zinc, bismuth and antimony, duli ensurels for enamelling iron vessels. These shades.

The composition of smalt varies considerably, the silica is usually from 56-70% potash 12-22%, cobait oxide 6-16%; it frequently contains small quantities of alumina, ferrous oxide, and lime. Smalt is a very permanent blue pigment largely used for the production of blue colours with vitreous substances It was formerly used for colouring starch, paper, etc., but for this and mony other purposes it is now almost superseded by artificial ultramarine, which,

however, is for less permanent.

Uses of Cobalt in the Ceramic Industry. -In the form of oxide, cobalt as largely used in the ceramic industry in three main ways (1) As a body stain, (2) as a glaze stain, and (3) as an underglaze or on glaze stain

(1) As a body stain cobalt is used because the clays used for nearly all pottery bodies contain iron which gives a yellow tinge to the fired goods The added cobalt counteracts this by pro-

ducing a complementary blue coloration which tralises the yellow and gives a pure white plour. For carrying out this process black oxide of cobalt is mixed with about 10% of china clay, thoroughly ground with nater, and passed through a 200 mesh sieve. The product is added to the pottery body in the wet state in the proportion of about 1 lh cobalt oxide to 1 ton of dry pottery body. The of blue, cobalt oxides in larger quantities are used. In combination with oxides of manganese, nickel, and chromium, all shades of blue and green may be obtained with cobalt oxide

(2) As a glaze stain the use of cobalt is most widely known in pottery manufacture because of the rich blue colours possible. The characteristic royal blue colour is obtained by adding 5% cobalt oxide to a glaze of high lead content. The oxide may be introduced either in the raw state or after fusing with a soft frit in s erucible, running into water, and grinding with the necessary quantity of glaze, Ali shades of blue, blue green, celadon glazes, etc., and mickel.

recovered. The mass is left at a high tempera- either smelt blue, which gives an intense

used, ENAMELS .- In the form of smalt robalt is used for the colouring of blue enamels. The oxide is also used in small quantities in the production of white enamels, thus neutralising any yellow due to the presence of iron. By the addition of cobalt oxide to certain raw mixtures and waste enamels a beautiful black enamel is produced which when added to other black enamels intensifies the colour; the rompounds such as sulcate, sluminate, and phosphate are largely used, as they give a more uniform colouring than the oxide, Cobalt oxides up to oxides are found to increase the adherence of

the enamels (T H. Gant, 1c).

Sulphides, - Coballous sulphide, CoS, 19 formed when aulphur is projected on to red hot cobalt, its formation being attended by incandescence and fusion The freezing point curve of the system Co-S has been determined sulphur by K. Friedrich up to 33 5% (Metallurgie, 1908, 5, 212) and indicates the formation of the compounds Co,S, Co,S, and CoS. The following sulphides have also been reported by versous workers; Co.S. Co.S., Co.S., CoS., end Co.S., Arsenides.—The compound CoAs, occur

in the mineral abutterudite in Norway, and CoAs, occurs in the important mineral smallule. The freezing point curve of the Co-As sense has indicated the existence of the compounds Co.As.; Co.As., Co.As., and Co.As (K. Friedrich, Mctallurge, 1908, 5, 150). The compounds Co.As., CoAs, and Co.As. have been prepared by Ducellies (Compt. rend. 1908,

147, 424). Antimonides .- The compounds CoSb and CoSb, here been indicated during the examination of the Co-Sb series by Ducelliez (Compt.

rend, 1903, 147, 1048).

Nitrides.-The nitride Co, N is described by Beilby and Henderson (J.C.S. 1901, 79, 1251); oxide to I ion of dry pottery body. The tho nitride Co.N. by Vournasos, Compt. rend. exact quantity necessary is determined by 1913, 188, 889; d. Bergstrom, J. Amer. Chem. trial. Sometimes the cohalt is added as Soc. 1924, 48, 2811). The axide Co.N.); chloride For colouring bodies various shades highly explosive (Wohler & Martin, Ber. 1917, 50, 586)

Silicides.-When metallic cobalt is hested to 1,200°-1,300° in elicon tetrafluoride vapour, the compound Co₂Si is formed as grey metallic crystals. This same compound, together with CoSI and CoSi, has been prepared in electric furnaces. Other silicides have been reported hy Jenge (Z. anorg, Chem. 1921, 118, 105).

CARBONYS.—There ere two carbonyls of cobalt, the tetracarbonyl, Co₄(CO), and the tricarbonyl, Co₅(CO). The tetracarbonyl is prepared as follows: cobalt oxists is heated. until completely converted into the oxide, which is then washed free from traces of chlorides may be made by mixing the cobalt oxide with the which hinder the formation of the carbonyl oxides of iron, chromium, manganese, copper, and dried at 120°. The oxide is placed in austable apporatus and reduced to cobalt by (3) For under glazes or on-glaze colour stains means of hydrogen at 300°, and a pressure of 5

atmospheres. After reduction is complete the hydrogen is displaced by earbon monoxide, the pressure being increased to from 40 to 250 atmospheres and the temperature reduced to 150°-250°, when varying quantities of carbonyl are collected in a tube attached to the outlet of the apparatus and immersed in a freezing mixture.

The tetraearbonyl forms orange-coloured crystals of sp.gr. 1.73 which melt at 51° and decompose above that temperature. It is isoluble in water but is soluble in most organic solvents and in nickel earbonyl. It is not attacked by aqueous acids, but concentrated acids and the halogens decompose it forming the corresponding salt, carbon monoxide, and hydrogen:

$Co_2(CO)_8 + 2H_2SO_4 = 2CoSO_4 + 8CO + 2H_2$

In contact with air, a deep violet-coloured compound is formed which is most probably a hydrated basic carbonate. The crystals are best preserved in a scaled tube in an atmosphere of carbon monoxide. When the tetracarbonyl is heated to 60° one quarter of the carbon monoxide is evolved at a regularly decreasing rate, leaving cobalt tricarbonyl, no intermediate product being formed. When heated to 100°, one-quarter of the carbon monoxide is evolved almost instantaneously, the rest coming off very irregularly and comparatively slowly (Mond, Hirtz, and Cowap, J.C.S. 1910, 97, 798).

Nitrie oxide reacts with cobalt tetracarbonyl slowly at ordinary temperature, but instantaneously at 40°, partly decomposing it into tricarbonyl and earbon monoxide and partly converting it into the nitroso-tricarbonyl Co(CO)₃NO, which is a cherry-red mobile liquid (Mond and Wallis, J.C.S. 1922, 121, 33).

Salts of Cobalt.—The soluble salts of cobalt are prepared by the solution of the oxide, carbonate or metal in the various acids. They are generally pink and deliquescent, forming pink solutions when dilute, but when dried or in concentrated solution they are blue. For this reason solutions of chloride or nitrate of cobalt are used as sympathetic inks, such writing being almost invisible until brought near a fire, when the letters appear blue. They gradually reals orb moisture and against become invisible. This change in colour has also here used for preparing "floral hydrometers" by tinting artificial flowers which in damp weather remain pink but in dry weather turn violet or blue.

When ammonia is added to solutions of cobalt salts oxygen is absorbed, giving rise under various conditions to many complex series of salts containing ammonia and known as cobalt-ammine salts which have received much atten-

tion (r. Coordination Compounds)

Cobaltons fluoride, CoF₂, is prepared by heating the chloride with ammonium fluoride in a current of hydrogen fluoride, and forms rose-coloured prisms slightly soluble in water (Poulene, Compt. rend. 1892, 114, 1426). The hydrated fluoride, CoF₂,2H₂O, is prepared by dissolving the oxide or carbonate in hydrofluoric acid; on evaporation rose-red crystals of the above composition are deposited (Edminster and Cooper, J. Amer. Chem. Soc. 1920, 43, 2419).

Cobaltie fluoride, CoF₃. has been prepared by O. Ruff (Z. angew. Chem. 1928, 41, 739) and by O. Ruff and E. Aseber (Z. anorg. Chem. 1929, 183, 193) by the action of fluorine on cobaltous chloride in a quartz tube at 150°. When heated to 250° in bydrogen it is reduced to cobaltous fluoride, the reaction being complete at a red heat.

Cobalious chloride, CoCl₂, is obtained by warming powdered metallie cobalt in chlorine gas; it ignites, forming blue crystalline seales of the anhydrous salt which can be sublimed in a current of chlorine. It is also prepared by dissolving the oxide or carbonate in hydrochloric acid and evaporating the solution. According to the temperature at which it crystallises, it may contain 6, 4, or 2 molecules of water of crystallisation, the last at the higher temperatures. The commercial salt contains 6 molecules of water and is violet in colour.

Coballous bromide, CoBr₂, is formed as a green fused mass of the anhydrous salt when bromine vapour is passed over metallic cobalt at a red heat. This salt absorbs water from the atmosphere, yielding a dark red liquid owing to the formation of CoBr₂6H₂O.

Cobaltous iodide, Col, is produced in the anhydrous state by heating metallic cobalt in iodine vapour. It is also obtained with evolution of heat when finely-divided cobalt is warmed

with iodine and water.

Coballous sulphate, CoSO₄,7H₂O, is prepared by the solution of the oxide or earbonate in dilute sulphurie acid. It erystallises in red crystals isomorphous with magnesium sulphate soluble in 24 parts of cold water. The following hydrates CoSO₄,6H₂O, CoSO₄,4H₂O and CoSO₄,H₂O are known, each of which on heating loses its water forming an infusible, opaque mass of a rose-red colour.

Cobaltie sulphate, Co₂(SO₄)₃,18H₂O, is obtained when a cooled acid solution of cobaltous sulphate is electrolysed in a divided cell. It is deposited in blue needles soluble in water, which solution rapidly decomposes into cobaltous sulphate. It has also been prepared electrolytically by S. Swanu, jun. and T. S. Xanthakos (J. Amer. Chem. Soc. 1931, 53, 400), who have shown that it has strong oxidising properties

towards many organic compounds.

Coballous nitrate, Co(NO₃)₂, in the anhydrous form, has been prepared by A. Guntz and F. Martin (Bull. Soc. ebim. 1909, [iv], 5, 1004) by the action of nitrie anhydride or of a solution of nitrie anhydride in nitrie acid on hydrated cobaltons nitrate and also by the action of finely-divided metallic cobalt on a solution of silver nitrate in liquid animonia. In this form it is a slightly pink powder which begins to decompose at 100°-105°. The hydrated nitrate, Co(NO₃)_{2.6}H₂O, is obtained by dissolving the oxide, hydroxide, or earbonate in dilute nitrie acid, when a carmine red solution is obtained and on evaporation this furnishes reddish-brown deliquescent erystals. Other hydrated nitrates containing nine and three molecules of water have been prepared. Cobalt nitrate solution is used as a reagent in blowpipe analysis.

USES OF COBALT SALTS IN PAINTS AND VARNISHES.—Certain cobalt salts, such as the cobalt arsenate, Co₂As₂O₃BH₂O. It found aretate, borste, resunate, cleate, inoleate, "color frequently as a peach-coloured incustation on resunates," bemoate, "timpate," and cobalt arsenued ores, being produced by their as hydroxide are used as driers in the manufacture of paints and varnishes. Organie salts of cobalt have been found to be the most convenient and efficacious driers and have the additional advantage of causing no discoloration of the paint. To some extent cobalt salts also enable the lower-priced semi drying oils such as soya, fish, and even cotton seed oils to be aubstatuted for the higher priced linseed oil in the preparation of paints. The driers in the liquid and solid forms are also used in the manufacture of waterproof fabrics, linoleum, oilcloth, patent-

leather, artificial leathers and similar industries, When the arctate is used for drying linseed oil, from 0.2 to 0.4% is necessary. The oil is beated at 150°-200° and the acetate is stirred in until dissolved. Soys oil is treated similarly

The borate is prepared by adding a solution of cobalt sulphate to a cold solution of borax; the precipitate formed is filtered washed with cold water, and dried,

The benzoate is prepared by adding cobalt carbonate to a solution of benzoic acid natil neutral; the filtered and dried product is

ground to powder.

For the preparetion of the resinate the best American water white rosin is used which should be completely soluble in turpentine and ether, have a melting point of 75°-85°, and a saponification value of 167-180. The resinate may be prepared either by the precipitation er the fusion method. For the precipitation method the resin, suspended in water, is heated to 100° and a 20% solution of sodium hydroxide is gradually added with stirring. A solution of sodium chloride is then added to the clear brown solution The precipitate of sodium resinate is dissolved in hot water and a 10% solution of cobaltous chloride is added to precipitate the sage green cobalt resinate which is washed and dried. The fused resinate is prepared by heating the resin to 140° and adding cobalt hydrate with constant stirring. The temperature is then raised to 170°-180° and the melt poured into pans and allowed to cool. The precipitated resinate contains about 7.5% cobalt and the fuscel resinata 4-4 23%.

The fatty acid driers are usually made with haseed oil, although other oils, such as soya, tung or walnut, are sometimes used. The linseed oil is completely saponified by gradoally adding 10% sodium hydroxida to the oil heated to 100°. The clear brown solution is then diluted with hot water and a solution of sodium chlorida added. The granular soap formed is removed, redissolved in water, and reprecipitated with brine. The purified soap is again dissolved in water and a 10% solution of cobaltous chlorida solution added. The precipitated cobolt "lineleate" is filtered, washed, and dried at 80°.

When 15% of a good drier is dissolved in haseed oil at a temperature of about 120° it should cause a thin film of this oil to dry at room temperature to a removable skin in 7-8 hours (cf. T. 11. Gant, I c.).

COBALT BLOOM. Erythrite, hydrated composition.

COBALT BLUE or THENARD'S BLUE v. COBALT, this vol. 218c.

COBALT BROWN, BRONZE, GREEN REO, YELLOW e. COBALT, this vol. p. 219c. COBALT-GLANCE, SULPHARSEN.

IDE OF COBALT P. COBALTITE.

COBALT PYRITES, LINNÆITE . CORALT., this vol. p 214b.

COBALT SPEISS E. COBALT, this vol.

2144. COBALTITE or COBALT-GLANCE. Sulpharsenida of cobalt, COAsS, containing Co 25-35%, with Fe 2-5%. It frequently occurs as well-developed crystals, fine specimens coming from Tunaberg and Hakansbods in Sweden and Skutterud in Norway; these are cubic with parallel hemihedrism and with the same characteristic form as crystals of the momorphous mineral iren-pyrites. Granular and compact masses also occur. The colour is tin-white with a reddish tinge and a brilliant metallic lustre ; the atreak is greyish black. Sp gr. 62, hardness 51. A bed of pure robaltite avrraging 0 2 m in thickness occurs at Dashkessan, Gort Teffes, Caucasus In the Cobalt area of Ontano massive cobaltite and small crystals are present with the more abundant amaltite. Under tha name sehis the mineral, found at Kherti, in Rapputana, is used by the Indian jewellers for producing a blue enamel on gold and ailver.

COBLAC: CELLULOSE PLASTICS COCAINE AND OTHER ALKALOIDS OF ERYTHROXYLUM SPECIES. The genus Erythroxylum (Fam. Erythroxylacen) is princ, pally found on the western aide of South America, although indigenous apecies occur in India, Africa and Australia Chiefly three kinds, derived originally from Bohvia and Peru, are cultivated

1. Erythroxylum coea.

2 E. coca, var. novogranglense (=E, carlhagense Jacq)

3 E truzillense Rusby (E. Spruceanum) On the London market four kinds of coal leaves are distinguished Husnuro (Bohvan, from 1), Truxillo (Peruvan, from 3); Java (frem 2), Ceylon (from 2 or 3).

The alkaloids obtained from coca leaves fall into four groups, viz :

The cocaines, which on hydrolysis yield ecgonine, methyl alrohol and benzoic, cinnamic,

or truxillic acids; Cocame: Methyl benzoylecgonine.

Cunnamoylrocaine, Methyl cinnamoylecgonine. a Truxilline (y-isatropy leocaine): Methyl atruxilloylergonine.

β-Truxilline (δ isatropyleocaine): Methyl βtruxilloy lecgonine.

2. The pseudotropeines are closely related to the trupemes and are hydrelysed into pecudo tropine and an acid.

Benzoyleegonine.

4. The hygrines, which are derived from pyrrolidine:

Hygrine and Cuscohygrine.

The two most important producing countries of coca are Bolivia and Java, but it is difficult to judge the world production of coca-leaves. In South America a considerable amount of erude eocaine is manufactured which does not come into the following statistics.

EXPORT OF COCA LEAVES (From the Annual Reports of the League of Nations (nos. in kgs.): 1933 1934 1932 1035 Bolivia 347049 335807 205011 170189 131955 Dutch Indies 209131 161591 104671 125155 117264

9 44598 44598 64598 49598Formosa 85542 76776 72374 89721 167896 Pern

Within the last years the coca production has much decreased, in 1924 Java produced as much as 1,071,000 kg. At the same time the number of coen plantations has decreased from 112 to 57 and the cultivated area from 2,112 to 864 hectares. Details of the production may be obtained from the Annual Roports of the League of Nations. The total alkaloid content of Javn leaves is higher than the total alkaloid in South American coca. In different years the yield of total alkaloids varies considerably (Java, 90% of 1908: 1-2.5%; 1909: 1.22%). total alkaloids present in South American coea is 1-cocnine, Java coca contains only 0.5% cocaine, the same percentage of cinnamyl-cocaine, and approximately 1% of truxillines, but these alkaloids can easily be transformed into l'ecomine. Coca-leaves experimentally grown in India contained 0.4-0.8% total alkaloids, chiefly cocaine (quoted by Henry, "Plant Alkaloids," 2nd ed., 1924, p. 96). The market for Java coea is Amsterdam; the London and Hamburg markets deal mostly with cocaleaves of other origin and with crudo cocaine.

The coca plant is a shrub which, when carefully cultivated, can be grown for about 20 years. The leaves are plucked in the early morning and must be dried and ground the same day; they are then packed and shipped in tin boxes, avoiding any access of moisture, which is deleterious to the nikhloid content (see Martell,

Pharm. Zentr. 1928, 69, 551).

For history of coca, botnnical details, cultiva-tion, etc., see Tschirch "Handbuch der Pharmakognosie" (Leipzig, 1923, vol. III, 1, pp. 309-326).

EXTRACTION OF COCA.—Either coca-leaves or crude cocaine of South American origin are extracted. The general outlines of the extractions are known, but many details, e.g. the use of solvents and the preparation of different coesine salts, are kept secret by the manufacturers.

The leaves are either extracted with dilute acids (see Ullmann, "Enzyklopadie der technischen Chemie," Berlin, 1929, vol. III, p. 453) or moistened with lime water or soda-solution plant material, see Klein and Sonnleithner

3. The acylegonines in which the carboxyl- and exhausted with Et2O or benzene (see group is free and only the alcoholic group is Duilius, Chem.-Ztg. 1930, 54, 31, or Schwyzer, esterified:

"Die Fabrikation der Alkaloide," Berlin, 1927, 85ff). Also extraction processes with H,SO4+EtOH as solvents (Squibb) and many other modifications are known.

(a) Extraction suitable for Java coca: The leaves are exhausted with dilute H2SO4; after eareful addition of soda the total alkaloids nre extracted with benzenc or tolucne. Extraction of these solvents with 10% H2SO4 yields a solution of the crudo total sulphates. Tho total alkaloids are hydrolysed with a small excess of dilute H₂SO₄ (boiling for 12-18 hours), when MeOH, benzoie, einnamie, truxillie acids, and eegonino are formed. The acids are removed and the mother-liquors concentrated; from the concentrated solution ecgonine is precipitated by EtOH. Thus all ecgonine present as cocaine, cinnamylcocaine, and the truxillines is available for the subsequent "synthesis" of cocaine. Ecgonine is first esterified with MeOH and H2SO4, and then benzoylated with benzoyl chloride or benzoic acid+POCl.

(b) Extraction suitable for South American coca: The leaves are moistened with limewater and extracted with Et2O. Dilute acctic neid extracts the total alkaloids from the ethereal solution. Crude cocaine is precipitated with soda-solution. This is dissolved in approx. 10% acetic acid, and at a temperature not exceeding +5° impurities are exidised with permanganate until the red colour persists for 30 minutes. The solution is then neutralised with numonia and the total alkaloids pre-cipitated with 5% soda-solution. The dried precipitate, which is brown in colour, is dissolved in Et.O and manganeso dioxido filtered off. The othereal solution is concentrated, when puro cocaino erystallises out. The mother-liquors are completely freed from Et.O and treated with light petroleum, which dissolves any cocaine still present. The residue consists largely of isatropylcocaine, which is hydrolysed and transformed into cocaine as mentioned above.

Analyses of Coca-leaves.—Tho official "Amsterdam-analysis" of coca-leaves is not published, the values obtained include cinnamyleocaine and truxillines. In London and in Hamburg the content of crude cocaine is given according to the methods of Harrison and of Gilbert. The estimation is based on the extraction of the leaves with n mixture of Et₂O, CHCl₃ and ammonia. The organic solvents nre then exhausted with dilute H2SO, and the basified solution re-extracted with Et.O. evaporated extract is dissolved in excess of N/10 H2SO4 and titrated back with N/50 KOH using cochincal or iodeosin as indicator (see "United States Pharmacopæia," 8th Rev.). Other descriptions use light petroleum as extraction medium; the further process is similar, the nlkaloids from the ethereal solution being dried to constant weight and then weighed. See also critical survey of Bierling, Pape and Vichöver (Arch. Pharm. 1910, 248, 303) and de Jong (Arch Pharm. 1911, 249, 209).

For microchemical identification of cocaine in

Official in the "British Pharmacoporia," 1932,

are: Cocaine and Cocaine HC1; official cocaine-containing prescriptions are: Lamella Cocaina and Lozenge of Krameriæ et Cocaina. Official in the "United States Pharmaeopœia,"

1936, are: Cocame and Cocame-HCL

Cocaine (I), C12H11O1N, [a] -164° (in CHCl,), m p. 98°, volatile above 90°, crystallises from EtOH in raonoclinic, four- to six-sided prisms It is easily soluble in EtOH (I part in I0 parts), Et,O (1:4), benzene, carbon disulphide, CHCl, (1:95), acctone and light petro feum, slightly soluble in H.O (1 600) saturated squeous solution is alkaline to himus. has a slightly bitter taste, and produces numbness when applied to the tongue.

(1) is precipitated from solutions of its salts by ammonia, caustic alkalis, and alkali carbonates When (1) is boiled with H₁O benzoyleegonine and MeOH are produced; hydrolysis with mineral acids and with caustic alkali yields l ecgonine (II) and d pseudoecgonine respectively, together with McOH and benzoie acid. Alka line hydrolysis is 1,500 times more rapid than

acid hydeolyais,

For dissociation constants of (1) and (11), see Kolthoff (Biochem Z. 1925, 162, 309-12). For test for identity and purity, see "British Pharmacopega," 1932, p. 124, and "United States Pharmacopega," 1936, p 123.

Cocaine Hydrochloride is the salt chiefly Cocaine Hydrochioride is the sait cheefy used in medicine It crystaltees from EIOH in short prime, [a]₀—71 95° (in 2% aqueous solution), in p. 200–202°. The sait is soluble in 0.5 parts of H₂O and in 3 parts of 90%. EIOH Test for purity ("Birtish Pharmacopera," 1932) to 8 millithres of a 2% aqueous solution add 0.3 millithres of N/1 H₂SO₂ and then 0.5 millithres of N/5O permanganate address volet colour is improduced. solution, a clear violet colour is produced, which in the absence of dust and at a temperature not exceeding 20° does not completely fade within 30 minutes (limit of reducing aubstances and of cunnamylrocaine) Similar test for purity in "United States Pharma-

Maclagan's Test. If a 10% ammonia solution is added to a solution of pure (I) HCI, a ery atalline precipitate of (I) must alowly be formed, a milky appearance indicates the presence of amorphous alkaloids, especially truxillines (for details of test and other tests, see "British Pharmacopous" and "United States Pharmaconceia"). Cocaine does not give characteristic colour-reactions; for Vitali's rolour-reaction, see Hardy (J. Pharm. Chim. 1921, [vii], 24, 3251

OTHER SALTS: (1) H.CrO4, H.O, mp 127, sparingly soluble in H.O, is produced when potassium chromate is added to an acid solution of (I) HCI Other sparingly soluble salts are B. H.PtCl. BHCl HgCl. BHl·l. (per-iodide), etc. (1) HCl forms aparingly soluble precipitates with Mayer's reagent, phosphomolybelic, phosphotungstic and alarotungstic acids (see Rosenthaler, "Der Nachweis organ ischer Verbindungen," 2nd ed., p. 732).

(Oesterr. Bot. Zeitschrift, 1927, 76, 263) and (I) HCl, see Heikel (J.C.S. 1909, 96, II, 104); Brandstetter (Chem. Zentr. 1922, IV, 729]. Nyman. Björksten (Z. anaj. Chem. 1915, 54 60. Nyman Björksten (Z. anai, Chem. 2018, 675); Garsed and Collie (J.C.S. 1901, 89, 675); Gersed and Collie (J. G. 1901, 89, 675); 1933, vol. IV, part 3/l, p. 540). (1) may be identified by the absorption spectrum, Bruster signified by the absorption appertum, Bruster (A. 1927, 91); Essenbrand (Amer, Chem. Abstr. 1926, 20, 2722); Castille and Ruppel (A. 1928, 924). Bontempi (Amer, Chem. Abstr. 1932, 26, 1862); Fischer (bid. 1933, 27, 4352) de Cocanne (de verasive) has been isolated from cora leaves by Luebermann and Greel (Ber 1890, 23, 509, 292). The silkalon is probably not natural but formed duming the extraction of the control of the con extraction. [a]_D +42 2° (in CHCl₃), mp 46°, gives a sparingly soluble nitrate.

Cinnamoylcocaine(111), C₁₈H₂₅O₄N, [a]₀
--4 7° (m CHCl₅), m p. 121°, crystallises from a mixture of benzene and light petroleum in resettes of needles. (III) HCl,2H,O, mp. 176 (dry), flattened needles from H,O, platinchlorade, mp. 217°, aurichlorade, mp 156° When (III) is warmed with HCI, it is by droly sed to I regonine, cionamic acid, and MeOH.

Truxillines (natropylcocames), Cas H45O4N4 Hesse (Ber. 1889, 22, 665) and Liebermann (Bet 1888, 21, 2342) isolated from Peruvian coca-leaves two amorphous alkaloids which were first named cocamine and isococamine,

a Truxilline (IV) (cocamine, y lastropylcocaine) · amorphous white powder, mp 80°, easily soluble except in light petrolcum,

β-Truxilline (V) (secocamine, 8 satropyl cocame). amorphous, ainters at 45° and decom poses at 120°. Both truxillines on seid hydrolyass yield ecgonine, MeOH, and a or β truxille acids (C,H,O,), respectively.

Benzoylecgonine, C, H, O,N, [a]0 -63 3' (in H,O); crystallises from water with 4H,O in needles, m p. 92° or 195° (dry). It is obtained by boding (1) in H.O. It was also obtained in small amounts from cocs leaves (see Skraup,

Monatah 1885, 6, 556).

C, H13O, N H,O, [a]0 ! Ecgonine (11), -45 4° (m H,O), m p. 198° (anhydrous), crystal bees from dry alcohol in monochnic prims with 1H₁O, easily soluble in H₁O, less in EtOH, insoluble in Et₁O, CHCl₁, and other organic solvents. (II) is the final basic hydro ly tie product obtained by action of acids on the cocames (see introduction); it is an amphotene aubstance and when esternied with MeOH and benzoylated, (1) is obtained. By variation of the acid chlorides and the alcohols, many homologues of (I) have been prepared.

d-Egonine (better d ψ-egonine) is dextro-cotatory, m p. 254° (261°); it erystallises from dry EtOH in tablets. d ψ-Egonine is formed when the cocames are bydrolysed by alkalis. d- Eegonine is not an optical but a stereoasomeride of (11). Lake the latter, it forms a series of exters; from d & ecgonine d cocaine (methy) benzoyl d & eegonine) has been prepared.

Tropacotaine (VI) (Benzoyl v tropeine) CasH18O2N, has been found in Java and Peruvian coca-leaves (Giesel and Liebermann, Ber 1891, 24, 2336, of also Reens, Chem. Zentr. 1920, 1V, 103). (VI) is optically inactive. ener verbindungen, "2nd ed. p. 732).

In p. 49°, it crystallises in needles, soluble in For quantitative estimation of (I) and EIOH and Et,O, insoluble in H,O. When

benzoic acid and 4-tropine (m.p. 108°, b.p. 240°) which is a stereoisomeride of tropine since it can be oxidised to tropinone. Tropine and ψ tropine are mutually convertible. Dissociation constants of (VI), see Kolthoff (Biochem. Z. 1925, 162, 307). Analytical differences between (I) and (VI) see Reichard (Pharm. Zentr. 1908, 49, 337). For synthesis of a ring-homologue of (VI), see Blount and Robinson (J.C.S. 1932, 1429). (VI), under the form of the hydrochloride, is the only other coca-alkaloid used in mediciae.

CONSTITUTION OF THE COCAINES .- The cocaines are esters of the amphoterie substance ecgoniae with MeOH and benzoic, cinnamic, or truxillic acid. Ecgonino and its derivatives differ throughout from tropine and its derivatives by CO₂. A derivative of ecgonine (anhydro ecgonine) was transformed into a derivative of tropine (tropidine) (Einhorn, Ber. 1890, 23, 1338); this fact proved that (II) is a tropine carboxylic acid. This is also confirmed by Willstätter's degradation of (II) to subcroae. The elucidation of the tropine-structure led Willstätter (Ber. 1898, 31, 1202, 2655) to propose the following structures for (II) and its derivatives:

(II): $R_1 = R_2 = H$. (I): $R_1 = C_6 H_5 \cdot CO$; $R_2 = Me$. (III): $R_1 = C_6 H_5 \cdot CH \cdot CH \cdot CO$; $R_2 = Me$. (IV) and (V): $R_1 = (C_9 H_8 O_2)_2$; $R_2 = Me$.

Formula (B) represents (VI). Wolfes and Hromatka (Amer. Chem. Abstr. 1935, 29, 4365) have isolated an interesting new tropane-derivative from coca-leaves for which they suggest the formula (C):

SYNTHESES OF COCAINE.—There are four stereoisomerides of ecgonine and cocaine possible, three of which have been obtained synthetically, r-Ecgonine and $r \cdot \psi$ -ecgonine have been resolved into d- and l- forms. These two forms different states of the second states of the s forms differ in the spatial arrangement of the hydroxyl-group (cis-trans-isomerism as in tropinc and stropine); the third form which has

heated with HCl or baryta it is hydrolysed to | discussion, see Willstätter and Bommer, Annalen,

1921, 422, 18).1. Willstätter and Bode (Ber. 1901, 34, 1457; Annalen, 1903, 326, 42) obtained sodium tropinone earboxylate by interaction of sodium tropinone with CO2 and sodium. This on reduction with sodium amalgam in weakly acidic solution gave a mixture of r- ψ -eegonine and r-eegonine (see additional note, Willstätter, Wolfes, and Mäder, Annalen, 1923, 434, 134).

2. A second synthesis of $r \cdot \psi$ ecgonine is described by Willstätter and Bonnaer (Anaalen, 1921, 422, 15): Diethyl succinyldiaectato is condensed with methylamine, the resulting pyrrolederivative is catalytically reduced to a pyrrolidinedicarboxylate which by a Dieckmann condensation is transformed into tropinoaccarboxylic ester. This ester is identical with the product of synthesis 1.

$$\begin{array}{c|c} CH_2-CO-CH_2-CO_2Et \\ \hline \\ CH_2-CO-CH_2-CO_2Et \\ \hline \\ CH=-C-CH_2\cdot CO_2Et \\ \hline \\ NMe \\ \hline \\ CH=-C-CH_2\cdot CO_2Et \\ \hline \\ H_2+Pt, followed \\ \hline \\ by Dieckmann \\ condensation. \end{array}$$

3. From the technical point the third synthesis of Willstätter, Wolfes, and Mader (l.c.) is the most valuable. Succinic dialdehyde, methylamine and the dipotassium salt of ethyl acctonediearboxylate are condensed to yield ethyl tropinonecarboxylate (cf. Robinson, J.C.S. 1917, 111, 762).

$$\begin{array}{c|ccccc} CH_2-CHO & CH_2-CO_2Et \\ & + & \\ + & \\ CH_2-CHO & CH_2-CO_2Et \\ \hline & CH_2-CH-CH-CO_2Et \\ \hline & \\ - & \\ & CH_2-CH-CH_2 \\ \hline \end{array}$$

Reduction of the last substance yielded r- ψ -ecgonine (1), r-ecgonine, (2) and the third racemie ecgoniae (cis-trans-isoaierie tropanol-carboxylie acid). On resolution (I) yielded d-aud l-ecgonine, the former, on esterification with MeOH and benzoylehloride gave d-cocaine (d-y-cocaine, see earlier). The bitartrate of d-v-cocaine is called psicaine, it is also used as local anasthetic and is a product of some importance. If (2) is resolved and esterified d- and l-cocaine are obtained, the latter is identical with natural cocaine.

Hygrines.—The hygrines were discovered not been resolved differs by the cis-trans-iso- by Lossen (Annalen, 1862, 121, 374) in Peruvian merism of its carbovyl-group. (For theoretical coca-leaves. Liebermann showed that this

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alkaloid fraction by distillation in ruces can be separated into the two alkaloids hygrine and cuscohy grine. Hygrine (VII), CaH15ON, [a]D-13°, bp.

92-94°/20 mm. or 193-195°/760 mm, is a strongly alkaline liquid, yielding a characteristic picrate, mp. 158°, and a crystaline owme, mp. 116-120°. By oxidation with CrO₂ hypra card (C,H₁₁O,N) is obtained which was synthesised by Wulstatter (Ber. 1900, 33, 1160) Annalen, 1903, 258, 91). r-Hygrine was synthesised by Hess (Ber. 1913, 46, 3113, 4104).

Cuecohygrine, C13H24ON, (VIII), is optically inactive, bp. 185°/32 mm, forms a crystalline oxime, m.p 53-54°, is miscible with H,O, a crystalline hydrate is known {m p. 40°), and the salts are crystalline The nitrate is characteristic, it crystallises from EtOH, m p. 200° (decomp). (VIII) is very labile; by influence of acids or alkalis it is easily trans formed into (VII); when oxidised with CrO, hygric acid (A methylpyrrolidine 2 earboxylie acid) is obtained. Various papers deal with the constitution of (VIII), but no final decision down covering the insect is preserved, but in the can be taken between the two possible alterna tives (see Hess and co workers, Ber 1915, 46, 1986, 1920, 63, 781, 1921, 54, 2310, Annalen, 1925, 441, 137).

PHYSIOLOGICAL ACTION OF THE COCAINE Chour .- Cocame is used in medicine as a local ancesthetic, it is mydriatic and highly toxic. It acts chiefly on the central nervous system, death occurs from paralysis of the respiratory centre. Tropacocaine is the only other cocacentre. Tropacocame is the biny out.

Alkaloid used in medicine For parcaine, see
Cottheb (Z. physiol. Chem 1923, 130, 374)

For details of cocame action, see Henry, "Plant Alkaloids," 2nd ed., p. 112, and text-books of deposits the carminic acid in a crystallios pharmacology.

COCAMINE r. COCAINE. (this vol p 224c). COCCERIC (COCCERINIC) ACID E.

Insects (r Countral Wax and Fat).

FAT AND WAX.

a COCCINIC ACIO, COCCININ . COCREMEAL COCCININ r. COCHINEAL, (this vol. p. 2001) COCCINONE v. COCHINEAL, (this vol. b.

2284). COCCULUS INDICUS OF INDIAN

BERRY C. ANAMIRTA PANICULATA.

COCHENILLIC ACID & COCHTYEAL Ithis vol. p. 2285)

COCHINEAL. This natural diestuff was used as a dye and cultivated by artificial means m its native country, Mexico, at a remote period of history. Cochineal consists of the dried body of an insect, the Coccus cacts, which lives upon a species of cartina (the Nopalea coccinellifera S.-Dyel or Nopal), a plant which is found in the wild condition, but which, for the sake of the msect, is cultivated in gardens which are termed Kopaleries. The insects are collected before the commencement of the rainy season, and are killed by immersion in boiling water. or are enclosed in a linen bag and placed in an oven; by the latter process the peculiar white former care is lost

In Mexico and Central America two varieties of cochineal are known-the home grown or fine cochineal (grana fina), and the wild or forest cochineal (grana silvestra) The former u more valuable than the latter, and is richer m colouring matter, After 1830 the cultivation of cochineal was introduced into Spain and the Canary Islands, Algeria and Java, but the most productive of these newer plantations were those of Java, Since the discovery of the coal-tar colours, the consumption of cochineal as a dyestuff has gradually decreased, and at the present time it is only employed in a minor degree According to Liebermann (Ber. 1885 18, 1969) cochineal contains about 10% of colouring matter.

Carminic acid, the colouring matter of cochineal, was first isolated by Pelletier and Caventou (Ann Chem. Phys 1818, [11], 8, 250) and was subsequently examined by many chemists. Schützenberger (ibid 1858, [iii], 54, 52) succeeded in obtaining carminic acid in a crystalline condition by precipitating the colouring matter as its lead compound, decomposing this with hydrogen sulphide, and crystallising the liberated earminic acid from alcoholether The most simple method of purification of carminic acid, however, is that devised by Miller and Rohde (Ber. 1897, 30, 1762) A solution of the crude colouring matter in five times its weight of water is diluted with four times its volume of acetic acid; the filtered liquid, on standing over sulphure scid, gradually condition.

Carminic acid crystallises in red prisms easily Schl. soluble in water and alcohol with a purple red colour. It possesses no melting point, but darkens at 130°, and at 250° becomes quite COCUMENT FIG. 300 Wig.

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C₂H₂₀O₁₃.
By the action of alcoholic potassium acctate, Perkin and Wilson (J.C.S. 1903, 83, 139) prepared a mono- and a di- potassium salt, whilst Liebermann, Höring and Wiedermann (l.c.) obtained hexabenzoylearminic acid as an orangecoloured powder, easily soluble in benzene. Octa-acetylearminic acid, golden-yellow needles, melting gradually at 155-165°, is readily prepared by the action of acctie anhydride in presence of zine ebloride or sulphuric acid (Miller and Rohde, I.c.).

Constitution of Carminic Acid.—By the prolonged action of boiling nitric acid on carminic neid, W. de la Rue obtained nitrococcussic acid, which was subsequently shown by Kostanecki and Niementowski (Ber. 1885, 18, 250) to be

identical with trinitrocresotinic acid:

When carminic acid, dissolved in 50% acetic acid, is treated with an excess of bromine, and the solution digested at the boiling heat, two substances, a and β -bromocarmines, are produced (Will and Leymann, Ber. 1885, 18, 3180).

a. Bromocarmine, C₁₀H₄Br₄O₃, colourless needles, m.p. 247-248° (decomp.), the more sparingly soluble substance, when oxidised with potassium permanganate in selution gives dibromomethylhydroxyaldchydobenzoic acid and dibromomethylhydroxyphthalic anhydride:

Miller and Rohdo (Ber. 1893, 26, 2603) observed that when a bromocarmino is treated with hot caustic soda solution it yields, in addition to a purple-red colouring matter, dibromomethyl-hydroxyphthalic acid and bromoform. As a result of this reaction, these authors assigned to a-bromocarmine the constitution of methylhydroxytetrabromodiketohydrindene (1), for Zincke (Ber. 1887, 20, 3227; 1888, 21, 2388) had previously shown that dibromodiketohydrindene (11) under similar treatment yields both phthalic acid and bromoform.

 β -Bromocarmine, C11H5Br5O4. crystallices

recently Dimroth has modified this expression to | (Will and Leymann) or 238° (Miller and Rohde), and is converted into a bromocarmine by the action of bromino in 50% acetic acid solution. As a result of their investigation, Miller and Rohde ascribed to \$\beta\$-bromocarmino the constitution of a metbyldibydroxytribromo-anaphthaquinono (III or IV). This suggestion

was supported by the fact that bromehydroxynaphthaquinono (V), on treatment with bromine and caustic soda solution, is converted into dibromodiketohydrindene (II) (Zineke, l.c.). Further support for this view was obtained by Rohdo and Dorfmüllor (Ber. 1910, 43, 1363), who found that by reduction with zine dust and acetic acid and subsequent acetylation, β -bromocarmine gives the compound (VI),

colourless needles, m.p. 208°. Simultaneous hydrolysis and oxidation converts this into tho substance (VII), orango prisms, m.p. 258°; and it thus appears that by the latter treatment an hydroxyl group has entered the quinone nucleus. The diacetyl derivative melts at 233°. When distilled with zine dust, this product, and also β-bromocarmine itself, give naphthalene.

VII.

Subsequently, Dimroth (Annalen, 1913, 399, 1) oxidised earminic acid with hydrogen per-oxide in aqueous caustic soda, using cobalt sulphate as a catalyst, and obtained 2:6. dihydroxy-8-methyl-a-naphthaquinone-3:5-dicarboxylic acid (VIII) which be orientated by comparing its colour reactions with those of synthetic 2:6-dihydroxy-a-naphthaquinone, and also by its conversion into carminazarin (XIV) by treatment with acid permanganate. When (VIII) is warmed with water, carbon dioxido from acctone in orange needles, m.p. 2322 is eliminated with the production of 2:6-

brominated in glacial acetic acid at 40°, yields the 7-monobrome- derivative which on treatment with hydrobromic acid gives a-bromo carmine, whilst with bromine in cold methyl sicohol the product is \$-bromocarmine, which thus has the structure (III) assigned to it by Miller and Robde.

IX.

Liebermann and Voswinckel (Ber 1897, 30, 688, 1731) studied the exidation of carminic acid with alkalino permanganate at the ordinary temperature, and in this way succeeded in producing two important acids, cochenilic (cochinelic or cochinilic) and a coccinic acida

Coch nilie acid, C10H4O2, colourless needles, m p. 224-225° with evolution of carbon dioxide, is tribasic and at 260° is converted into methyl hydroxyphthalic anhydrido (X) When heated with water in a sealed tube at 210° it yields cresotime acid (XI), and in the same manner at a lower temperature, 170°, gives a coccinic acid

(XII), the second product of the oxidation of

By oxidising carminic acid with potassium permanganate at 0° in the presence of sulphune the other hand, alkaline oxidation by means acid, Dimroth (Ber. 1909, 42, 1811) obtained of hydrogen peroxide below 20° gives use to carminazaria, garnet red needles, decomposing cochenilic acid. Dimroth thus formulated at 240-250°. It possesses the constitution coccinin and coccinion as derivatives of an-

duhydroxy-8-methyl-a-naphthaquiuoue-5-ear (XIV) and is very similar to isonaphthaann boxylic acid (IX). This compound, when (XV). Its alkaline solutiou, when treated with a stream of oxygen, is quickly decoloursed with formation of 56 dicarboxy-4-hydroxy-o-tolyl-

glyoxyhe acid (XVI). A further point of resemblance of carminazarin to isonaphthazarin is shown by its behaviour with mitric seid in glacial acetic acid, for whereas the latter cives

tetraketotetrahydronaphthalene, tho former the analogous carminatarinquisons (XVII) which, when heated with water or scetie acid, passes back to carminazann. azarinquinone is converted into a diphenana, C24H14O2N4, hy treatment with an alcoholic solution of o-phenylenediamine.

XVIL

Coccenin, according to Hlasiwetz and Grabowski (Annalen, 1867, 141, 329), is produced when carminic acid is fused with caustic potash. It crystallises from alcohol in straw-yellow needles or leaflets, dissolves in alkalis with a yellow colour which, by air oxidation, develops first a green, then violet, and, finally, a purpls tint. Dimroth (Annalen, 1913, 399, 1) examined this Johnson (Annalen, 1913, 399, 1) examines us product in detail, preparing it by fruing examines and with eaustic potash at 170-2007. To it be pale yellow crystalline ictra-acetyl, derivative, in p. 212-244. When occurrent was outless of year, or oxygen, in dilute acdium hydroxide solution, the colour changes above destroyed occurred, and when the pure violet colour had been obtained, actification wideld occurred. (XII), the second product of the Unisatum us, earning each a -Coccinic each, C₂H₂O₂ colour less needles, mp. 230°, proved to be identically with m-bydroxyuritie and (Dippenheim and Plaff, Rer. 1874, 7, 920). It thus follows that the continual (XIII).

Cocumon in reconstruction and product of the continual con reduction with zine dust and ammonia; on

Dimroth further concluded that carminie acid itself is a derivative of anthraquinone, for by distillation with zinc dust in an atmosphere of hydrogen, a 5% yield of hydrocarbons of tho anthracene series was obtained. After oxidation of the mixture of hydroenrbons he isolated anthraquinone, and possibly a-methylanthraquinoue. Further, by treatment of earminie acid with boiling dilute sulphurie acid, a 10% yield of trihydroxymethylanthraquinone carboxylie acid was obtained, and this acid, when heated with water at 230-240°, yielded tri-hydroxymethylanthraquinone by loss of earbon dioxide. Dimroth therefore formulated earminie acid:

An examination of kermesic acid, the colouring matter of kermes, by Dimroth and Fick (Annalen, 1916, 411, 315) showed that this acid is a derivative of hydroxyauthrapurpurin. The similarity of the absorption spectra and dyeing properties of kermesic acid to those of carminic acid led Dimroth and Kammerer (Ber. 1920, 53. [B], 471) to infer that the latter is also a derivative of hydroxyanthrapurpurin.

Under suitable conditions carminic acid is reduced by rine dust and acetic acid to a leuco-compound which is converted by atmorpheric oxidation into deoxycarminic acid,

thranol and anthraquinone respectively, the | carminic acid differs from carminic acid only in production of cochenillic acid indicating the positions of the earboxyl and one hydroxyl and loss of the β -hydroxyl group in the purpurin one methyl group: carminic acid contains two hydroxyl groups in the 1:4-position is shown by its oxidation to a diquinone, from which carminic acid is regenerated by the action of acetic anhydride and sulphuric acid. Parallel reactions are observed in the case of kermesic acid and hydroxyanthrapurpurin.

When carminic acid is warmed with acetic anhydride in the absence of a catalyst, a hexaacetyl derivative is formed, and two hydroxyl groups in a positions in the nucleus remain unnectylated, since the substanco is readily oxidised to an unstable diquinone, reduced by sulphurous acid to the parent substance. With acetic anhydride and sulphuric acid at the ordinary temperature, earminic acid yields an octa-acetyl derivative. Since only four hydroxyl groups are present in the anthraquinone nucleus, it follows that the remaining four must be present in the side-chain, to which tho composition C₆H₁₃O₅ has been previously assigned (Dimroth, Annalen, 1913, 399, 13). This side-chain can only be a saturated univalent hydrocarbon radiele with five oxygen atoms, four only of which have been shown to be present as hydroxyl groups; hence it is coneluded that the remaining oxygen atom is present in the ethercal, aldehydic, or ketonic form, and its formula must then be modified to C₆H₁₁O₅. Accordingly, the empirical formula for carminie acid becomes $C_{22}H_{20}O_{13}$, and an examination of recorded analyses showed that they agree as well with this as with the older formula. Hence carminie acid is:

$$C_{22}H_{20}O_{13} = HO_{2}C \parallel OH C_{6}H_{11}O_{6}$$

Carminie acid is optically active and has [a] 645 +51.6° in aqueous solution This phenomenon cannot be attributed to the structure of the nucleus, since kermesic acid, and also the less elosely related laccaic acid, are optically inactive; it therefore appears that the side-chain contains at least one asymmetric carbon atom.

According to Miyagawa (Mem. Coll. Eng. Kyushu Imp. Univ. 1926, 4, 99), carminic acid, when treated with ozone in aqueous solution, yields an optically active product from which, by oxidation, an optically active acid, C6H12O6. is obtained. On reduction with sodium amalgam this acid yields a sugar, C6H12O5, which is not identical with any of the known methylpentoses. The side-chain in carminic acid has thus the probable structure CH₃·[CH·OH]₄·CO—.

Technical Preparations of Cochineal—Am-

moniacal Cochineal, or carminamide, is prepared by allowing a solution of carminic acid in yellowish-red needles, the acetyl-derivative of ammonia to stand for some time; the clear liquid which melts at 245-250° (decomp.). Deoxy- is decanted and evaporated with, or without, addition of gelatinous alumina. The product | acid, has been proved to be a mixture of 15. appears to consist of carminic acid in which one Leto n-dotriacontanoic acid. of the hydroxyls has been replaced by an amino-group. Ammoniacal cochineal dyes much bluer

shades than cochineal itself, and was sometimes used in conjunction with the latter; it also found some application in the blueing of bleached

Dyeing Properties of Cochineal.-Cochineal has been little employed in cotton dyems, but was very largely used in aik and wool dyeing. It has now been practically replaced

by the azo scarlets.

A crimson shade is produced on wool by mordanting with aluminium sulphate and eream of tartar and dyeing in a separate bath with cochineal, whilst a very fiery acarlet is obtained by the employment of stannous and sometimes stannie chlorides For scarlet, wool can be mordanted with atannous chloride and tartar, and dyed in a separate bath with cochineal. A single bath method, using atannous chloride, oxalic acid and cochineal, has, however, been very largely used for this purpose. On wool mordanted with chromium or iron, cochineal gives purple, slate or lilso colours, but these mordants are not employed in practice.

Cochineal red on wool possesses considerable fastness to light, but has the defect that weak alkalis and soap cause it to acquire a duller or

more bluish shade.

Silk is dyed a good crimson shade by mordant ing with alum and dyeing with an extract of cochineal. In scarlet dyeing, silk is preferably first dyed yellow, then mordanted with "nitro-munate of tin," and finally dyed in a second bath with the assistance of cream of tartar. Silk can also be dyed in a single bath. For cochineal indicator r Vol II, p. 6394. A. G. P. and E. J. C.

COCHINEAL FAT AND WAX. extracting cochineal insects (Coccus cach) with boiling benzene (Liebermann, Ber. 1885, 18, 1975; Liebermann and Bergams, soid. 1887. 20, 959) or chloroform (M Becker, Brochem. Z. 1931, 239, 235) a fatty material, which can be separated into an alcohol- and ether insoluble wax, cocceen (c. 05-1% of the doed meeta) and the ether soluble cochineal fat (c. 7-14% of the insects). The fat, as examined by Huerre (J. Pharm Chim. 1911, 4, 56) had an sodine value of 50 and an acid value of 89; it contained only 3% of unsaponifiable matter and yielded 8% of glycerol. The fatty acids appeared to consist of a mixture of myristic, oleie and linolie

Coccerin (m p. 99-103°C.) (cocceryl coccerate) was shown by Liebermann (l.c.) to be an ester of cocceryl alcohol with coccerrance acid. Cocceryl alcohol, originally believed to be a Cas (Liebermann) or Cas (Becker) dihydrie alcohol, has been identified by Chihnall and coworkers (Biochem J. 1934, 28, 313) as 15-keto-n. tetratriacontanol,

Correspondingly, cocceric acid, which was in 1931-1932 to 679,160 tons in 1934-1935

CH3 [CH2]18 CO [CH2]11 COOH, and n-triacontanore acid, CH, [CH,], COOH

(Chibnall, Lc., cf. sbid. 2189, 2209).

COCINIC ACID. A term now obsolete apphed by Saint-Evre (Ann. chim. 1847, 20, 99) to an acid, in p. 34 7°C., anpposed to be undecrie acid, separated from coconut oil Like "umbellake acid," which was isolated from the seed fat of the Californian bay tree (Umbellularia californica Nutt , Fam. Lauracen) by Stillmann and O'Neili (Amer. Chem. J. 1883, 4, 206) and also believed to be an undeeylic acid, "cocinic acid" appears to have been, in fact, an impure pre-paration of lauric acid (dodecylie acid). Unde cylic acid (" cocime acid ") is also absent from chaulmoogra oil, in which its presence was reported by Moss (Pharm, J. 1879, 39, 251)

COCLAURINE is an Important alkaloid, as at is the mother-substance of the whole group of the "biscoclaurine-alkaloids" (cf. Kondo and Tomita, Arch. Pharm. 1936, 274, 65).

It was isolated from Cocculus laurifolius by H Kondo and T. Kondo (J. Pharm Soc. Japan, 1925, No. 524,876). It crystallises in colouries plates, m.p. 221°, [a]_D -17 01°, easily soluble in hot EtOH and acetone, sparingly so in cold EtOH, acetone, Et,O, CHCl, insoluble in

benzene and light petroleum.

B.HCI, needles, m p 264°; methiodide prums. mp 155° (tri acetyl and tri-benzo) com-pounds). By oxidation of the methine base, Kondo obtained oxalic, anisic, and a derivative of phthalic acid (Chem. Zentr. 1927, I, 2203) By ethylation of the phenolic O'H groups, H. Kondo and T. Kondo obtained degradation products which demonstrated conclusively the following atructure (J. pr. Chem. 1930, [ii] 125.

For absorption spectrum of coclaurme, Kondo, Chem. Zentr, 1929, 11, 752, 1012.

COCOA. The carao tree is an evergreen shruh indigenous to Mexico and the tropical regions of Central America, whence it was introduced into Europe by Cortes in 1528. It belongs to the family Sterculacese and comprise several species, that most highly valued and cultivated being the Theorema Cacao, so called by Lannæus, from βεός (god) and βρώμα (food) in allusion to the esteem in which the beverage prepared from its seed was held by the natives of Central America, and to the Mexican name of the tree-Cacouatl or Quacahault.

PRODUCTION .- The world production of cocos bas ateadily increased, rising from 577,600 tons formerly thought to be a C11 or C12 bydroxy. Of the latter total, the Gold Coast produced

259,400 tons and Brazil 98,700 tons. Other | vated. The pods are smaller whilst the beans producing countries are Caracas, Central America, Ecuador, Essiquibo, Grenada, Guayaquil, Peru, Trinidad, Venezuela and the West Indies, Ceylon, East Indies, Philippines, Mauritins, Madagasear, San Thome, Cameroons and the northern parts of Australia. The exportation of eocoa is of increasing importance in the trade of certain portions of the British Empire. Over 60% of the world trade is of imperial origin, whilst about 95% of the raw coton used in the United Kingdom comes from the Empire. The quantity of raw cocoa retained for home consumption in this country has increased from nearly 50,000 tons in 1926-1927 to over 70,000 tons in 1935-1936. largest importing country in 1934 was the l'uited States of America with 197,000 tons, Germany being second with nearly 100,000

CULTIVATION.—The caeao tree thrives only in tropical temperatures, in districts which are moist and sheltered from winds. Thus, those places where it is cultivated with success lie in latitudes between 25° north and south of the equator and at elevations seldom exceeding 1,000 feet. The soil affording best results is well drained, alluvial sandy loain. McDonald (liap. Coll. Trin. 3rd Ann. Rep. 1933, p. 36) gives data relating to suitable and unsuitable soils in which the organic matter in good specimens varies between 2.8 and 4.0% whilst the available P2O5 is between 63 and 149 parts per million. Manuring is not often necessary although stable manure can be applied with advantage, and to sandy soils the addition of superphosphate in the proportion of 1-3 lb. per tree promotes fertility. Potassium fertilisers tend to increase the weight rather than the number of the beans. The tree usually grows to a height of about 25 ft., although occasionally it reaches 40 ft., but under cultivation it rarely exceeds 18 ft. It begins to bear fruit when 3 or 4 years old, attains full productivity in a further 4 or 5 years, and may be expected to bear for 40 or 50 years. Fruit is produced all the year round, but harvesting usually takes place only twice a year-from February to April and again in October and November. This requires great skill and judgment, as there is danger of the fruit being cut off too early or the flowers of the later crops being danaged. The fruit, which is suspended on a short thick stalk, consists of an elongated pod, varying in shape between a melou and a enember, measuring from 5 to 12 in, in length and 2-4 in, in diameter. Iuside the pods are beans, varying in number from 20 to 40, and embedded in a rosecoloured mucilaginous pulp. The trees have been divided into three main classes, depending | upon the fruit and other botanical differences: Criollo, Forastero, and Calabacillo. The first class, in which may be included those of Java and Ceylon, produces the finest type of beans, but the trees are more difficult to cultivate and

are flat and have a more astringent taste than the other classes. When fresh the beans of all classes are usually white, although occasionally ranging through red to purple, but after drying and exposure to air and light they acquire a vellow, red, or brown colour and become hard, brittle, and less bitter.

FERMENTATION .- After removal from the pods the beans have to be freed from the adhering pulp. This is occasionally done by artificial or sun drying and subsequent rubbing and sorting. This method, however, produces a bitter product, quite unfit for the manufacture of chocolate or high-class cocon. The usual procedure includes fermentation, which, whilst facilitating the removal of the pulp, improves the flavour and aroma of the final product. To promote fermentation the beans were formerly buried in the ground or in trenches covered with plantain leaves. Owing, however, to the adherence of earthy matter to the beans and the difficulty of control and handling, this process is not now generally employed. The beans are now placed in casks or boxes or heaped on the floors of sheds. The fermentation process usually lasts from 2 to 7 days, the temperature in that period rising to about 48°C. Care has to be taken to maintain an even temperature throughout the mass and this is achieved by shovelling the heaps periodieally or transferring from one container to another. The process of fermentation is first alcoholie, owing to the action of a yeast fungus Saccharomyces Theobroma Preyer, and later acetie, set up by Mycoderma aceti. F. L. Stevens (Bull. Dept. Agric. Trinidad and Tobago, 1925, 21, Part 1, 27-35) expresses the opinion, based upon extensive investigation, that successful curing depends upon (1) non-development of acidity through avoidance of undic exposure to bacteria and yeasts; (2) complete killing of the embryo by heat; and (3) maintaining the oxidising enzymes in an uninjured condition. A product of superior colour and aroma is obtained by keeping the beans at a temperature of 40°-60° for several days out of contact with bacteria and yeasts. During fermentation a large proportion of the pulp is gradually transformed into a liquid ("sweatings"), which is allowed to drain away. When the beans are satisfactorily cleared of the pulp and have assumed the requisite aroma and colour they are removed to sieves or troughs and thoroughly washed with water if such a eleaning process be considered necessary. In order to improve the quality of the beans Defren (U.S.P. 1750795) suggests that they should be soaked in successive changes of water for a sufficient time to remove at least the greater proportion of the soluble bitter ingredients. As an alternative to the fermentation process, a method has been devised whereby the beans are treated with a 1% solution of sodium earbonate for 10 minutes at 45°. It is are less regular in their yield of fruit. The wall stated that cocon prepared in this way is perof the pod is soft and thinner than the other preformed sugar. Theobromine is not present includes some of the best products of Central America. Calabacello is the most easily culti-

by fermentation or other means the beans are dred to a moisture content of 5-8% by heating sartificially or by subjection to the rays of the sun. Care should be taken to control the temperature, as above 50°C, the flavour is bleft. The hiss is form a low priced product sometime. to be adversely affected. If more than 8% of used, after grinding, in cheaper grades of cocos moisture is allowed to remain, the beans may become mouldy. In order to basten the drying of "cocoa tea." being infused as a better-finely powdered dired earth is sometimes apread which has the flavour of weak cocoa. They are over and intimately mixed with the beans. The mass is then raked, band rubbed, or "danced" with the bare feet, the friction thus set up also polishing the beans, thus improving their and Herrmann (Landw. Versuchs Stat. 1933, appearance, and removing mildew, thereby eliminating the possibility of the growth of mould during subsequent storage.

MANUFACTURE.-The preliminary processes of manufacture are :

(a) Sifting out of sand dust and small atones. (b) Picking out by hand of the empty beans and foreign matter, such as larger atones, grass, and wood.

(c) Roasting. (d) Husking hy breaking up the beans and separating the busks.

Berhardt (Chem - Ztg 1889, 13, 32) has found, from the results of actual factory working, that from the resures of actuar narroy working, that the losses in these operations amount to 2-3% in sifting, nearly 1% in picking, about 5% in roasting, and 12 or 13% in busking, making a total loss of over 20% Machinery is now in use which combines the first and second operations with the sorting of the beans into various sizes. The objects of cossting are to develop to the full extent the characteristic acoma and flavour of the cocos, to render the starch granules more soluble, to modify the tannin and other astringent ingredients, to render the and other assument ingrements to state the husby brittle and easily remosable, and to drive moisture from the nib, thus rendering it fit for the aubsequent operations. The process, which requires considerable care and experience, is conducted in cotary, cylinders heated to a temperature varying from 160 to 315°C, by coke fire, gas flames, or hot air or superheated steam in pipes hining the cylinders, Uniformity of coasting is facilitated by grading the beans so that they are approximately of the same size in an operation. Defren (Food Ind 1930, 2, 248), and in the manufacture of chocolate confechowever, objects to this usual method of roasting alleging loss of flavour, charring, and difficulty of control. He auggests that the arematic flavours are better retained by the to form an inducement for its abstraction from use of a stream of heated air to reduce the mois ture content to about 3%. The dampers should then be closed and coasting at about 100° con tinued After reasting, the beans are cooled and then gently crushed in a "kibbling" mill to fracture the crisp husk and break down the kernel into its natoral angular fragments or "nits," after which the nibs, husks, and hard rod shaped germa are separated by amnowing be prepared either by treatment of the cocoa and sieving. Care must be taken that the proand sieving. Care must be taken that the pro-ponder with steam or amnonia or akal such at

and chocolate and occasionally, under the name also used as fertilisers, and as an ingredient in eattle feeding materials. They have also been suggested for use as cattle bedding, but Mach 117, 265) express the opinion that they are not suitable owing to their low water absorption capacity compared with that of straw, whilst, if the preportion of theobremine is over about 100 their consumption by the animals would be dangerous

The nibs constitute the important part of the bean and form the basis of all coces and chocolate preparations, their ultimate use governing the method of treatment to which governing the method of treatment to wants they are subjected. This, however, does not apply to the first stage, in which the mis-are pulvensed in a hot mill. They are fed through a bopper on to revolving stones escared an a jacket beated to about 45°C. Here the nibs am speedily reduced to a molten mass owing to the high content of low-melting fat (about 30°C.). When the mass is considered to be completely melted and homogeocous it is allowed to run into moulds whem it cools and sobdifies It is then in a convenient form for atorage, transport, or further treatment, and it is in this state that much cocos forms an article of commerce under the title " block cocoa.

The form in which cocos to be consumed as a beverage is prepared for sals is known com mercially as "cocoa powder." The high peoportion of fat in block cocoa would render it indigestable and prevent complete mixture with water About 60 or 70% of the fat is therefore removed by subjecting the block coros to s pressure of about 1,200 or 1,400 lb per sq mat a temperature of 70°-80°C. The cocca butter flows out through a special onfice and s afterwards solidified in moulds ready for use tionery Although in this process the fat of "cocoa butter" was at first regarded as a waite product, its commercial value is now so high as the cocoa andependently of the demand for the defatted cocoa. The cocoa mass, which after this treatment contains about 25% of fat, is then cooled, erushed in a mill, and sifted, the resultant powder being cocoa from which the beverage is prepared. A feature of the cocos trade for many years has been the preparation and sale of so-called soluble rocoas. powder with steam or ammonia or ammonium will become brittle and break up into ampliparticles which may be lost in the busk. The shorts continue, on an exerge, about 120-125, of the weight of the bean, varying from 7 or 8% or the weight of the bean, varying from 7 or 8% or the middle beans of Ceylon and dava which weight of the the middle beans of Ceylon and dava who boiling water for the preparation of the beverace.

regarded as adulteration, but if other substances are added the fact should be disclosed. Thus it is a common practice to offer for sale preparations in which starch or sugar or both have been added to the cocoa powder. The designation of these by some such term as Composition of Roasted Nib.—This has "chocolate powder" is generally considered to been the subject of considerable investigation, be adequate indication of the presence of sub-stances other than cocoa. The addition of Chemistry and Manufacture," London, 1912) starch or sugar, if properly disclosed, is not to gives the following as typical results:

an emulsion is formed which retains the powder | be condemned. Indeed, it may well be claimed in suspension. These admixtures are not that the presence of the starch is of dietetic advantage, increasing the earbohydrate content and rendering the mixture more easily assimilated by those who find the pure cocoa product, with its high fat content, indigestible. (For preparations of cocoa, see Chocolate.)

	1.	2.	3,	4.	5.	6.
Moisture Fat Albuminoids Collulose Ash Gum Alkaloids Cacao red Pentosans Nitrogen Fibre Astringent matter Cane sugar Starch Cold water extract	5.23 50.44 13.26 6.40 2.75 2.17 0.84 2.20 — 6.71 4.20	6-3-8-5 46-9-52-1 11-6-21-1 3-3-6-6 2-9-4-8 0-3-0-5	3·7-4·4 45·3-54·4 7·4-13·0 2·4-3·9 ————————————————————————————————————	3·00 50·00 — 3·07 — — 2·50 2·80 — 11·60	5·86 50·30 — 4·05 3·87 — — 1·36 — 9·97 —	3·11 54·37 — 3·41 — — — — — — —

J. Bell, "The Chemistry of Foods," 1887, p: 76.
 Zipperer, "Untersuch, über Kacaoets," pp. 56, 57
 Helsch, Analyst, 1875, 1, 142.

N. P. Booth, Analyst, 1909, 34, 144.
 R. Adaa, Internat. Cong. App. Chem. VIIIC, 203.
 F. Bordas, Internat. Cong. App. Chem. VIIIC, 188.

gredient of the cocoa bean is the fat which, as already indicated, is, in itself, a valuable commercial commodity sold under the names "Cocon Butter" or "Cacao Butter." Its principal uses are for medical purposes and for addition to block cocon in the manufacture of checolate confectionery. Under the title "Oleum Theobromatis" the "British Pharmacopæia" describes cocon butter as "the solid fat expressed from the roasted seeds of Theobroma Cacao. A yellowish-white solid fat; odour elight, agreeable and resembling that of cocoa; taste bland and characteristic. Somewhat than stearic were present. brittle but softens at 25°C." Cocoa butter is

Cocoa Butter.—The most important in- and boiling alcohol from which it crystallises out almost completely on cooling. Lewkowitsch (J.S.C.I. 1933, 52, 236) gives the following as its approximate composition: a palmitooleo-stearin 55%, oleodistearin 25%, β-palmitodiolein about 20%. Lüning and Drudo (Z. Unters. Lebensm. 1931, 61, 491) tested several samples by Graf's method of fractional precipitation of the magnesium salts (Arch. Pharm. 1888, 228, 843) with the modification that the magnesium salts were dried at 95° till of constant weight. They also determined the melting-points of tho fatty acids and concluded that no acids higher

The following are some of the values of eocoa readily soluble in ether, chloroform, turpentine, | butter as published by various investigators :

	1.	2.	3.	4.
Specific gravity	0·990-0 998 (15°C.)	0-964-0 974 (15°C.)	0·8823-0·8829 (60°C,)	0.8831(60°C.)
	()	(12 0.)	0·8572-0·8581 (90°C.)	0-8562(99-9°C.)
Saponification value .	192-198	192-195		_
Iodine value	31-10	32-42	34-9-40-1	_
Refraction [n]	1-4565-1-4570	_	1-4568-1-4570	
Melting-point °C Redwood Viscosity	32-31	30–34	30-2-31-2	32-6
at 60°C.	-	_	99-1-101-3	102

¹ Elelon, "Edible Oils and Fats."
2 Whymper, "Cocoa and Chocolate."

Tate and Pooley, Analyst, 1921, 46, 229
 Knapp, Moss, and Melley, Analyst, 1927, 52,

from cocoa by the ordinary commercial method as high as 80 on an arbitrary scale. of expressing and that obtained by the method of extraction with a solvent as often followed in the laboratory may be of importance, par-ticularly when considering the possibility of the presence of some foreign fat Visser (Chem. Weekhiad, 1932, 29, 40) considers that the term "cacao butter" without further designstion should be applied only to the fat obtained by expression. Fincke (Bull. Off. Intern. Fabr. Choc. Carao, 1932, 2, 327) atates that when fat reason for the extraction of the fat it is some is extracted from chocolate for examination, ether and not light petroleum should be the solvent, as the latter will not dissolve out the wax which is likely to be present in the adulterant fat. Castiglioni (Ann. Falsif. 1935, 28, 24) givra two methods of differentiation between extracted and expressed cocoa fat. In the first the melted fat is shaken with glacial acctic acid and allowed to at and The fat on rising to the top is then examined by ultra-vinlet light and ened oils have the advantages of more uniform if extracted shows a hright yellowish-green fluorescence. In the second the fat is heated and elimination of "sugar bloom." Williams with alcohol, hydrochloric acid and antipyrine, (Chem. Trade J 1935, 95, 4) states that peanut when a pink colour indicates that the fat has and cottonseed oils hardened to 35° have a greater been extracted. Neither reaction is given by degree of contraction on cooling to 10"-12" (Chem. Umschau Fette, Orle, Wachse, Harre, because of the higher priportion of unsaturated 1931, 33, 265) found that whilst the refracto- glycerides in the former. Coconut oil when used is meter readings showed no marked difference readily detected by the various constants. Thus:

The differences, if any, between fat obtained the Haber-Lowe interferometer gave differences

Although, in the preparation of cocos as beverage, the removal of a considerable portion of the fat is desirable, the high commercial value of cocos butter, compared with that of the ray bean or the various preparations of cocos, has promoted the removal of the fat from the nib to an excessive degree and its replacement in chocolate confectionery by other fats of recetable origin. In addition to the economic times claimed that, for certain purposes, the admixture of a foreign fat is advantageous facilitating manufacture and providing a more stable product. The two fats most commonly nsed for this perpose are coconut or palm kernel stearin and the so called " illipe hutter," a name given in the chocolate trade to Borneo tallow. Recently hardesed coconut, palm kernel, peanut, and cottonseed oils have been used. The hardcomposition, better control of the melting point

	Sp gr IS S €	Saponifi- cation value,	lodine value	Ж.р. ° С.	Refrac- tion, [n]p	Der	Titre (f p of fatty stids, "C.)
Cocoa butter (Elsdon) ,	0 990- 0 998	192-198	31-10	32-34	1-4565- 1-4570	0.5	49-50
Coconut oil— Fryer and Weston Bolton	0 926	255-260 255-258	8–9 79–88	23-26 23-26	1-4486- 1-4492	1 6-1-0	21-25

Illipé hetter, however, does not display any [and Pooley (le) determined the following auch marked differences from cocos butter. Tate values :

		Sp gr co°C	Redwood Viscouty at 60°C	М р °С.	Jodine Value	M p of fatty acids.	Refraction,	
	Cocoa hutter	0 8823- 0 8829	99 0-101 3	30 2-31 2	34 9-40 1	47·7°- 49 2°	1 4568- 1-4570	
i	Illipé hutter	0 8823- 0-8840	100 7- 105-7		27-4-33 4		1 4561- 1-4573	

It will be seen that the respective values for average 3,150. Those obtained for a similar the two fats are so close that none affords by st. number of samples of illipé butter ranged from self an effective means of differentiation. Tale 3,800 to 4,771, average 4,60. Knopp, Most, and Robey proposed to multiply together certain and Melley (Analyst, 1927, 25, 452) coulder this of the constants determined in a standard method useful if applied with distriction but manner, thus obtaining a composite factor for each of the fats. The constants treated in this way were the specific gravity at 60°C, the sp gr. at 99°C., the viscosity, the melting point, the melting-point of the free fatty acids, and consider the titro "alue the most useful angle the reciprocal of the todice value. The comfactor which can be determined. That for posite factors thus obtained on twelve samples cocos hutter varies only between 49°C, and 50°Cof cocoa butter varied fram 2,839 to 3,347, whilst the mean value for Borneo tallow is

they suggrat that each worker should determine his own average constants. They further suggest the use of the "titre" value, particularly if other substitutes are suspected. They also

reputed cocoa butter is adulterated if it gives a "titro" over 50°C. A method proposed by Bywaters, Magss and Pool (ibid. 1927, 52, 324) takes advantage of the extreme supercooling to which cocoa butter can be subjected. It is based upon the "turbidity temperature" of the melted fat. The method appears to afford a valuable approximation to the percentage of illipé butter present. Pichard (Compt. rend. 1923, 176, 1224) also uses the minimum temperature of supercooling as a means of detection of adulteration. By plotting the temperature of a fused fat as a function of time, cocoa hutter from different sources gave the same curve, differing from those for all possible adulterants. The method suggested by Kochler (ibid. 1924, 178, 940) is based upon a comparison, preferably at 15-20°, of the number of drops of ethyl acctoacctato which must be added to a 20% solution in chloroform of the sample to obtain constant turbidity unchanged by the addition of another drop. The "turbidity number" varies considerably with temperature, and the sample and pure cocon butter should be treated simul-taneously under identical conditions. Hanus and Komorousová (Chem. Listy, 1925, 19, 391) define the "cthyl ester number" as the number of e.c. of N/10 potash necessary to saponify the ethyl ester in the distillate obtained from 5 g. of fat after conversion of the glyccrides into ethyl esters. The "numbers" for eoconut oil and cocoa butter are given as 23 and 3 respectively, referred to 1 g. of fat. Ashmore (Analyst, 1934, 59, 515) has brought forward an ingenious apparatus for observing the temperature at which molten cocoa butter crystallises. A beam of light led by internal reflection along a glass rod is projected through a small tube of the molten fat suitably housed in a darkened chamber. The appearance of the tube is noted during cooling. As soon as crystals of solid glyceride separate the tube is at once illumined by the Tyndal effect produced. The temperature at which this occurs can be noted with considerable precision and may be used for detecting the presence of certain adulterants. Thus genuine cocoa butter has a crystallising temperature of 20°C., whilst a temperature of 31.25 was recorded for Borneo tallow.

The variation in the melting-point of cocoa butter and adulterant fats due to varying time and conditions of crystallisation has been recognised by many workers. Fincke (Z. angew. Chem. 1925, 38, 572) recommends that the fat should be cooled slowly and with stirring. He found that for cocoa butter the temperature after 0.66, 2.5, and 14 days respectively was 32.3°, 32.3°, and 32.6°. Sabalitschka (ibid. 1925, 38, 1013) considers that, in order to obtain complete solidification, it is necessary to leave the melting-point tube on ice for several weeks before making the determination. In order to overcome the difficulty Tate and Pooley (Lc.) determined the melting point by placing

54.6°C., and they are of opinion that a sample of | very small flame and the temperature taken when the first signs of transparency were observed in the shaving.

THEOBROMINE (v. CAFFEINE AND THE ALKA-LOIDS OF TEA, COFFEE AND COCOA) is the principal alkaloid in eocoa and is closely related to caffeine which is also present to a small extent in eocoa. The proportion in which theobromine has been found to be present has varied considerably, probably in consequence not only of variations in the different type of bean but also, in some measure, of the different methods of estimation employed. The caffeine is usually included with the theobromine, from which, however, it can readily be separated by means of cold benzene in which caffeine is soluble but theobromine practically insoluble. According to Kreutz (Z. Nahr. Genussm. 1909, 17, 526) the proportions found have often represented only the free theobromine. Ho maintains that part of the alkaloid is combined and is not extracted until the cocoa has been subjected to hydrolysis. He found proportions varying from 2.8 to 5.1%. Wadsworth (Analyst, 1921, 46, 32) suggests a method of determination in which 10 g. of the fat-free material are triturated in a small porcelain dish with magnesium oxide and water, the mass is then refluxed with tetrachlorethane and, after filtering, the residue is again refluxed with the same solvent and the united washings distilled to small bulk. Precipitation is then effected with ether and tho precipitate dried and weighed. Wadsworth further states (Analyst 1922, 47, 152) that the theobromine content varies with the type of bean from 2.2 to 3.8% on the dry fat-free residue. Goryainowa (Khem. Farm. Prom. 1932, 6, 227) suggests a method embodying some slight modifications of that proposed by Wadsworth. During the process of fermentation the theobromine content of the nib decreases, whilst that of the shell increases. According to Churchman (J.S.C.I. 1926, 45, 149) the percentage in eocoa shell varies from 0.2 to 3%

ALBUMINOIDS.—These have been but little studied, but Stutzer (Z. angew. Chem. 1891, 4, 368) has classified the nitrogenous ingredients of

eocoa as follows:

(1) Non-proteids, substances soluble in neutral water solution in presence of copper hydroxide (theobromine, ammonia, and amino compounds).

(2) Digestible albumen, insoluble in neutral water solution in presence of copper hydroxide, but soluble when treated successively with gastric juice and

alkaline panereas extract.
(3) Insoluble and indigestible nitrogenous

compounds.

According to Stutzer's analyses, the proportion of albuminoids in cocoa powder manufactured without chemicals is between 17 and 18%, of which 10% is soluble or digestible albumen, the remainder (over 40% of the whole) consisting of insoluble and indigestible substances. Forster (Hyg. Rundschau, 1900, 314), however, maina shaving of fat upon mercury contained in a tains that the proportion of digestible albumi-porcelain crucible in which the thermometer noids reaches as high as 80%, his experiments was suspended with the bulb fully immersed. being made on the human subject, whilst Stutzer's The crucible was placed on a wire gauze over a were conducted under laboratory conditions.

STARCH, as in cleaginous seeds generally, is | MINERAL MATTER.—The ash in cocca nile not very abundant. Owing, however, to the occasional necessity to identify and estimate to Zippere being 3 %; in raw and 3 %; in the Approximately half the attempts of the control of the and proportion in the bean are important. The (which is rich in potash and phosphates) is starch grains are nearly spherical with a very indistinct nucleus. They have a tendency to the average when starch or sugar has been unite in small groups of 3 or 4 and rapidly lose added, and higher when fat has been abstracted the colour imparted by rodine. The average or when the cocoa has been adulterated with size is about 0.005 mm. The proportion in which it occurs in the nib has been variously estimated as from 2 to 2000, but many of the results are too high, owing, no doubt, to the conversion of cellulose into sugar when the acid method for the starch conversion is employed The diastase method is preferable in the presence of cellulose. The average amount of starch is from 4 to 500, or about 800 calculated on the

fat-free bean, TANNIN AND COLOURING MATTER-It has long been known that the raw bean, which is nearly white, contains an astringent principle resembling tannin which is rapidly oxidised to form "cocoa red," thus imparting to cocoa ita characteristic colour. Bell states that this astringent anbetance is different from the tannin of tea and coffee, and that its rapid change to cocoa red, even during the process of analysis, renders its quantitative estimation difficult. Adam (Analyst, 1928, 53, 369) found that the unfermented cocoa bean contains (a) a subatance belonging to the estechin series, (b) a catechu tannin, and (c) a compound of cocoa tannin and caffeine. The colouring matter which is aubsequently developed consists of rocoa red and cocoa brown, which are complex alteration products of the estechin and tannin originally present. The cocoa catechin undergoes alteration during the process of fermentation and is absent from the completely fermented bean. Fincke (Z Unters Lebensm 1928, 55, 559) states that both the cocoa red and the cocoa brown are tannin substances, the former occurring in variable amount in the fresh seeds, whilst the latter is gradually produced during fermentation, roasting and grinding. Steinmann (Z. Unters. Lebensm. 1933, 65, 454), on the other hand, states that the formation of cocoa red in the bean depends mainly upon the type of cocca; white unfermented beans contain no cocoa red which is found only in beans with violet cotyledons. Unfermented beans dried in the dark contain no cocoa red, but those aubjected to the action of light contain free cocos red which, however, disappears on fermentation. The increase in colour intensity on long fermentation is ascribed to the liberation of combined cocoa red, indured by the acids, chiefly acetic, formed during fermentation. This cocos, from that of mb, being 2-3 times as great, and red goes over slowly into cocos hown. The development of the dark brown red colour in the shell of the bean on drying is probably a photochemical reaction. Steinmann further states that whilst cocoa red is easily soluble in alcohol, cocoa brown will dissolve only with difficulty, but does so readily in an Sci. Pharmacol. 1920, 27, 355) gives, as a reason

soluble in water. The ash is naturally lower than husks or coloured with pigments, A high alkalimity in the ash of cocoa powders indicates treatment with alkali as dracribed in the preparation of "solnble" cocoa (see von Fellenberg, Mrtt. Lebensm. Hyg 1932, 23, 29). Cocoa Shell.—The chemical detection and

approximate determination of thell used in cocoa products is of use only in cases of gross adulteration. Microscopie methods are generally admitted to be more suitable. In this connection von Fellenberg (1bid. 1921, 12, 301) found that a asmple of cocoa powder contained stone cells 3-6 times the diameter of the cells characteristic of cocoa shell and resembling those of walnuts. These show mostly a thick wall in the interior of which are elongated pores. Sometimes no thick wall is noticeable, in which case the pores reach the penphery. They are notired in many cocoas rich in shell They are characteristic of the cocoa fruit shell and samples showing them contain shell of the fruit as well as that of the ared Their presence may therefore be taken as a certain indiration of adulteration with added shell. Koperberg (Chem. Weekblad, 1926, 23, 646) suggests that the stone cells in the defatted material should be counted and compared with standards containing a known percentage of shell. He expresses the opinion, however, that the result can be regarded only as an approximation, as commercial samples are not sufficiently uniform. Griebel and Sonntag (Z. Uniters. Lebenam. 1926, 51, 185) agree with this but suggest that it may be possible to set limiting figures for each rocce roduct based upon this method. Alpers (ibd. 1927, 54, 462) further draws attention to the fact that stone cells in cocoa products are deformed and broken up hy grinding and regard thould be had to this fact when romparing samples with standards. Plücker and Steinruck filed. 1931, 62, 361) suggest a methol fit cleaning the tissue, colouring with safranne and counting the cells. Winkler (J. Assoc. Of. Agric. Chem. 1935, 18, 427) states that crude cellulose is the most promising characteristic for indicating the proportion of shell in cocos products, the spread in percentage of this ronstatuent in mib and shell being much greater than that of crude fibre. The ash of shells differs

being richer in alkaline carbonates and silica. COCOA FEEDING MEAL.-The use of the produets of the cocoa bean in the preparation of feeding stuffs for animals renders important the findings of investigators who have expressed opinions adverse to such use. Rothéa (Bull. aqueous alkaline solution giving a deep for the unsuitability of coops shell as a hore brown colour. On hydrolysis cocca red de- food, the high percentage of theoremies, of composes into glucose, tannin and a resumens which be found 1-09%. He is of the opinion substance.

that uniterated cocca shell is unfit for animal that uniterated cocca shell is unfit of animal than the contract of the contr food. Marchadier and Goujon (Le Mans, J. Pharm. Chim. 1919, 20, 209) state that cocon shell deprived of its extractive matter 19 devoid of nutritive value, and its sale as food for animals should be prohibited. They agree with Rothia, however, in suggesting that the alkaloidal bases of cocoa shell are dangerously toric, and that the shell should be used only in small daily doses. With regard to cows, Bunger and Lamprecht (Milch. Zentr. 1926, 55, 49) find a decided increase in total protein of the milk when cocoa meal is fed, the casein increasing more than the albumin. Lactose, on the other hand, decreases somewhat. Aplin and Ellenberger (Vermont Agrie, Exp. Sta. Bull. 1927, 272) found that coeoa meal in the rations of milking cows increased the proportion of fat but reduced the milk yield so that the gain In fat production was small. They further state that the theobronine was responsible for theso changes, the easterne content producing no noticeable effect. Aplin (ibid. 1927, 271) also states that eocoa meal 19 not a suitable ingredient of hog or pig ration, causing litters of undersized pigs when fed to pregnant sows and everting a toxic effect on young pigs. Robinson (Olno Agric, Exp. Sta. Special Circular, 1932, 26) states that cocon bean is not of great value in fattening hogs, and attributes the poor results to the presence of theobromine and caffeine.

COCONUT, COCOANUT. The fruit of the cocoanut palm, Cocos nucifera L. The fibrous outer husk of the cocoanut yields the coir fibre largely used in the manufacture of matting and cordage. The dried fleshy endo sperm of the nut is the copra of commerce, or when specially cleaned, dried, and shredded forms the "desicented cocoanut" or "cokernut" used in confectionery. The press cake from oil extraction, known as "cocoanut cake," is a much valued eattle food.

Of the whole nut 30-50% consists of husk; the remainder, the eccoanut usually met with, comprises shell 27-28%, fleshy kernel 55-56%,

" milk " 17-18%.

The following are average analyses of cocoanut products:

	VIIIk.	Kernel	Desterated cocoanut	Coconnut	Copra.
Water	91.5	463	3.5	107	58
Protein	0.4	4 I	63	19 1	89
Fat	1.5	37.3	57.1	11.1	67.0
Carboliy drate	46	79	١	f 41·0	124
Fibre	0.0	3 1	31.5	141	4.1
Ash	0.8	1.0	1.3	4 0	1.8
	1 Ir	om ripe	nuts.		

The protein of coconnut flesh consists almost caurely of a globulin, conglutin (Ritthusen) or phytovitellin (Chittenden and Setchell) containing 17 8-18 4% N and 1 8% ash. A small amount of a proteine is also present. Johns, Finks and Gersdorff (J. Biol Chem 1919, 37, 149, also Jones and Johns, ibid 1920, 44, 283, 291) record the mitrogen distribution of the globulin as:

				0%	of total	N.
Amide-N .					7.99	
Humin-N .					1.5	
Cystine-N .					0.96	
Arginine-N.					29 5	Ł
Histidine-N					3 68	į,
Lysine-N .					641	y
Ammo-N of	filtro	ite			45 44	
Non-amino-I	V of	filtı	ate		46	

The composition of the products of hydrolysis

						%
Alanine						4.11
Valino .						3 57
Leueine						596
Serine .						1.76
Cystine						1.44
Aspartie ac	nd					5-12
Glutamie a	eid					19 I
Tyrosine						3.18
Plienylalan	ine					205
Proline .						554
Try ptopha:	n				p	resent
Arginine						15 92
Lysine						5 80
Histidine						242
Leucylvalu	ne a	nhy	dri	de		0 64
Ammonia						1.57

The milk of the cocoanut consists of an emulsion of oil in water stabilised by protein, the colloidally dispersed proteins being adsorbed at the oil-water interface. The aqueous phase contains dissolved sugar (Clemente and Villacorte, Univ. Philippines Nat. Appl. Sci. Bull. 1933, 3, 7). In the milk of young nuts sucrose is the principal sugar, and small amounts of glucose and mannitol are reported. The proportion of glucose and sucroso appears to vary considerably in the different stages of maturation (Dunstan, Trop. Agric. Mag. Ceylon Agric. Soc. 1906, 26, 377; Lahille, Bull. Econ. Indochine, 1920, 23, 1). According to Gonzalez y Sioco (Philippine Agric. Forest, 1914, 3, 25) the ripening is characterised by three stages: (1) commencement of endosperm formation when invert sugar and amino acids accumulate in the milk, (2) drying out of tissue of green nut when sucrose is produced, (3) oil formation in the endosperm during which the total solids of the milk diminish. In dried copra meal, Caray (Philippino Agric. 1924, 13, 229) found the following percentages of earbohy drate materials: sucrose 14 3, raffinose 2 4, galactose 2 4, pentoses 24, fructoso 1.2, glucose 1.2, cellulose 155, pentosans 22, starch 09, dextrin 06, galactan

Buchofen records the following mineral analyses:

analyses.				
-	Husk.	Shell.	Kernel.	Milk.
Total ash	1 63	0.29	0 79	0.38
K.O	30 71	45 01	5106	34.57
Na ₂ O .	27.56	$23\ 67$	2.66	13 96
CaÖ	4-14	626	3.10	7.43
MgO	2.19	1.32	1.98	3 97
Fe ₂ O ₃ .	054	1.39	0.59	trace
P.O	1 92	4 64	20 33	5 68
so,	3 13	5 73	8 79	3 94
SiO	8 22	4 64	1 31	2 95
CI	27.88	9 44	9 25	35 52

COCONUT,

Soc. Sci. Hyg. aliment, 1928, 16, 457j. A. G. Po.

fats [often known as the" nnt oils," or in German Elacis guineensis, and the aquara palm, the peras "Leimfette"), which are derived from the carp or fruit-flesh aurrounding the seed is sign seeds of various tropical species of the family rich in oil, which differs, however, considerably Palme, and which differ from other vegetable from the corresponding seed-fat; these pala fats by having high saponification values (235-265) in conjunction with high Reichert-Meisal Palit Oil. values (5-9) and Pnlenske values (10-20): the iodine values are fairly law (c. 8-25). The kernels of the cocount palm, Cocos nucifers L. fairly high Reichert-Meissl values, consequent which grows m all coasts and islands in the upon the presence of the acluble volatile fatty tropics. The tree is an posed by De Candolle ecids, caproic and caprylic acids, bring these fats. (cf. A. W. Hill, Nature, 1929, 124, 133, 151) into anperficial relationship with the mammalian milk fats; they differ essentially from the latter, however, not only in containing vegetable phytosterols (in the unsaponifisble fraction) in but also in that the nut oils do not contain Ceylon and Africa, so that it is now found any butyric acid. The presence of the volatile caprylic and capric acids differentiate the nut mis from other fats (such as those of the Dika Fat Group, q.v., and Laurel Fat Group), which likewise possess high saponification values but do not contain appreciable amounts of acids of lower molecular weight than lauric acid, and hence display very low Reschert Meissl and Polenske values.

Two fats of this group, tiz coconut oil and palm-kernel oil, are of very great commercial importance, whilst the trade in bebassu nuts and murumnru nuts, which has become established within the last twenty years, seems likely to mcrease. A number of units of this group, such as cohune unts (from Orbignya cohune Mart Dahlgren, v. infra), mocaya, Paraguay or gru-gru palm nuts (from Aerocomia selerocorps Math, treum and course must (from Mathematics, treum and storage must defeat and attracting must return Matt and Attracting must be must be must be must be defeated from portion of the Capitals exports desired for 1916, 44, 293, Bolton and Hewer, told the must fit be must fit be must be 35, 210), and the nuts in the societie or linais paim, Maximiliana regia Mart. (cf. Bull. Imp. Inst. 1916, 14, 8; 1927, 25, 1) necasimally appear on the market as sources of coconutlike fats Cocos bonetii, Becc. is being cultivated experimentally at Suchum in the Black Sea region; as might be expected, the nil from this species, which grows on the northern bant of distribution of tho genus Cocos, has a higher rodine value (erz. 23-24) than the fats from the tropical members nf the group, and it is therefore proposed to hydrogenate the nil, in order to render it more acceptable as a substitute for cocount ml (cf. S. Ivanov and Alissova, Chem. Umschau, 1929, 36, 401). The fats from a number 1929, 38, 401). The fats from a number of of their species, such as the hunty and must palms (Mourius 29.), the jassary palm (Astro-caryum pauray Mart), the purruma palms (Mourius 29.), the jassary palm (Astro-caryum pauray Mart), the purruma palm (Cocos 2907013 Drude), and uther aprecess of Atlanta, Marmiliana, etc., are utilized by the natures of the countries in which they grow for eduble purposes, as huming mils, or in pharmacy or soap-making; it is possible, also, 37, for the British Empire, Fullippica liked, as Build East Indier repetitive).

Small proportions of zinc occur in the endosperm that such fats may be included at times in the but not in the milk (Bertrand and Benzon, Bull. "coconut oil" prepared in such districts coconut oil" prepared in such districts. It may be pointed out that the fats discussed

in this article are ell derived from the sede COCONUT OIL GROUP. Under this (kernels) if the palms; in some instances, for heading is grouped a number in closely similar example, in the case of the common oil palm fruit fats will be discussed under the heading

Coconut Oil is the fat derived from the to have been indigenous in the Malayan Archipelagn, whence the sceds, well protected by the enveloping thick coir, were carried by sea currents eastward to the Pscific Islands and the place of the cholesterol of the ammal fats, coast of Central America, and to the west to growing wild, or under semi cultivation, on all teopical coasts in both hemispheres; it is bowever, especially abundant on the Malabar and Cocomandel coasts of India, in Ceylon, and in the East Indian end Pacific Islands. Apart from self-planted or "wild" coconst land, the extent of which cannot be calculated, at has been estimated (1930-1932) that about 7 million acrea are devoted to the cultivation of the coconut, of which some 96-97% ha in Aus and Oceania, and about 53% within the British Empire. Whilst the Indian acreage of some 1,330,000 acres is the largest area in any angle country, the whole of the crop is now consumed locally, leaving no balance for export, and the chief countries exporting copra or coconut oil, or both, are the Philippine Islands, the Dutch East Indies, the South Sea Islands, Ceylon, and 65% of oil) from the principal exporting countries are in the order of I to I million tons per annum, of which, from 1924-1930, 35% (on the average) was furnished by the British Empire (including Ceylon), and about 30% each by the Philippine Islands and Dutch East Indies. These shipments represent shout 57% of the total production, the halance of 43% being consumed incally by the copra producing countries. (For detailed studies of the etstistical and economic position of coronut oil and copra, from which the above figures have been extracted, see "Survey of Oilseeds and Vegetable Oils," No 2; "Coconut Palm Products," Empre Marketing Board, E.M.B. 61, Dec. 1932, H.M.S.O.; "Vegetable Oils and Oilseeds,"

Snodgrass, "Copra and Coconut Oil," Leland- kernels into boiling water and skimming off the Stanford Univ., 1928).

The coconut industry in Ceylon and Malaya is described by F. C. Cooke (Dept. Agric. S.S. and F.M.S. Gen. Ser. 1932, No. 8), who also makes a comparative report on the Philippine Islands industry in a later publication (ibid. 1936, No. 23; cf. Gothwaite, "Trade in Philippine Copra and Coconut Oil," U.S. Dept. Commerce, Trade Promot. Ser. No. 11, 1925; Snodgrass, l.c.).

The eccount palm flowers when about 5-7 years old, and begins to bear fruit from the age of about 8 years up to 30 or 60 years or more, according to the variety of the plant and cultural conditions. The large, drupaceous fruit consists of an outer skin and thick fibrous pericarp (from which the well-known "coir" is obtained) surrounding the stone or "nut," which consists of a hard woody shell (endocarp) within which is the coconut kernel or seed. The fleshy endosperm or "meat" which contains the fat is built up gradually from a milky juice as the fruit ripens. When the nut is mature, the milky juice has almost disappeared, and the endosperm forms the well-known hollow kernel, in which the remainder of the juice (the eccount "milk") is enclosed. The fresh "meat" contains from 30-10% of fat and 50% of moisture.

Since the earliest times in the history of man, coconut kernels have been used as articles of food by the natives of the South Sea Islands and of India: the oil first became known in Europe in the eighteenth century, but neither the oil nor copra was imported in any notable quantities until 1820 (cf. Soames, B.P. 5842, 1829). In the United States, which is now the largest consumer of coconut oil outside the copraproducing areas (apparent consumption, c. 260,000 to 330,000 tons per annum), the utilization of coconut oil in soap-making only dates from the beginning of this century, whilst its employment in the manufacture of margarine has only developed since the World War.

France new follows the United States as the second largest non-tropical consumer of the oil (c. 120,000 tons per annum), whilst Germany, which was by far the biggest consumer in pre-War days and held second place from 1924 to 1929, fell to third place in 1930, and was just superseded in apparent consumption by the United Kingdom with 92,000 tons in 1931.

The most primitive native method of obtaining the oil consisted in merely chopping up the kernels and exposing the pieces to the sun until the oil exuded and could be collected. In ladia, where at an early date the oil was extensively used for food and in pharmacy, more refined methods, which form the basis of present-day commercial practice, were employed.

After removing the fibre the nuts were split, the "milk" poured off, and the half-nuts exposed to the sun until the kernels were dried and could easily be detached from the shells; the dried kernels were then triturated and expressed in order to recover the oil. The residual mass ("coconut poonac") was used as a cattle-food. The best quality of oil—the original "Cochin oil" from the Malabar coast—was prepared by throwing the pounded, dried the trade as a designation of quality in copra, without especial reference to the mode of drying. In the somewhat primitive kilns employed, the fumes from the fire obtain ready access to the copra itself and impart to it a peculiar empyreumatic odour and also cause a certain amount of discoloration, and, occasionally, scorehing; the quality of the product (which is judged by the appearance of the copra and by the colour, etc., of the oil obtain-

kernels into boiling water and skimming off the liberated oil. The reputation of Cochin oil was such that the term has come to be used to designate the finest, whitest grades of oil, regardless of the place of origin or of the method employed to prepare the oil (as stated above, Indian-produced oil (and copra) is no longer exported to Europe). "Ceylon oil" was prepared in Ceylon by similar methods, but, owing to various circumstances, represented as a rule a slightly inferior grade of oil having a somewhat higher content of free fatty acid than the Malabar product.

A further expansion in the coconut industry followed when it was recognized that, whereas the fresh, moist kernels readily putrefy, causing hydrolysis and raneidification of the oil, welldried kernels (containing less than 8% of water) could be preserved for a considerable time without damage to the contained oil, so that the drying treatment developed into a manufacturing process for producing the dried kernel -copra-for export to Europe and the United States, where the oil could be expressed on the largo sealo with modern machinery. Formerly, apart from the Cochin and Ceylon oils (amounting to about 10% of the total), all the industrial eoconut oil (" copra oil ") was thus prepared in the consuming countries from imported copra, and it is still true to-day that copra bulks far larger in the exports from the producing areas than does coconut oil. There has been a growing tendency, however, within recent years to establish modern oil mills in the principal copraproducing districts, such as the Philippino Islands, Ceylon, Dutch East Indies, and Malaya, and to export the oil as such, so that in the period 1927-1930 about one-quarter of the aggregate exports of copra and coconut oil from the copra-producing areas was shipped in the form of oil (E.M.B. 61, l.c.). Formerly a sharp distinction existed between "sun-dried" copra and "kiln-dried" or "smoke-dried" copra; for a given typo of coconut, a "sun-dried" product, prepared by exposing the split nuts to the sun for several days, is the better, since it yields a paler and less acid oil; but in regions where the climate does not permit of this process, recourso must be had to artificial drying, and the split nuts are "kiln-dried" or 'smoke-dried" by sprending them on (bamboo) grids supported over a fire of coconut shells and husks (cf. F. C. Cooke, l.c., also Malay, Agric. J. 1936, 24, 167, 332; 1937, 25, 93). Even in Ceylon (which produces the highest-priced copra) it is now the usual practice to allow only a preliminary drying in the sun (for 1-2 days) and to complete the drying in the hot kilns (4-6 days); hence the term "sun-dried" has come to lose its original narrow significance, and is used in the trade as a designation of quality in copra, without especial reference to the mode of drying. In the somewhat primitive kilns employed, the fumes from the fire obtain ready access to the copra itself and impart to it a peculiar empyreumatic odour and also cause a certain amount of discoloration, and, occasionally, scorehing; the quality of the product (which is judged by the appearance of the respectively.

TABLE I. USCAL ANALYTICAL CHARACTERISTICS OF FATS OF THE COCONUT OIL GROUP.

	Coconut oil.1	Palm ketnel oil.	Cohune nut olL	Babsssu fat.	Murumura fat.
Melting-point	23°-28°C. (2>30°C.)	23°-30°C. (\$30°C)	18°-20°C.	22°-26°C.	32°-35°C.
Specific gravity	c. 0 874/22	c. 0 873/15	0 868-0 871/160	c. 0 867/100	
Iodine value ,	8-10 (7 5-9 5)	15-23 ³ (14-19)	9-14	8-17	11-13
Saponification value .	250-264 (<255)	243-250 (242-252)	251-256	246-263	237-242
Reichert-Meissl value .	6-8	5-75	68-85	5-9	c. 3
Polenske value	15-20	10-12	125-155	4-13 3 (nsually 11-13)	£.7
Bertram A-value	27 4-30 5	16 5-17(a)	l –	19 5, 24-1 (6)	87(c)
Bertram B-value Hehner value (insoluble	25-31	1 8-2 1(a)	-	1-9-20(b)	14(c)
fatty acids)	88-91%	c 90%	1	88-91%	c. 93%
Unsaponifiable matter.	c. 0 2% (>0.8%)	02-08%	0.5%	03-08%	03-06%
Titre (f p. of fatty acids)	20°-23°C.	20°-25°C.	c. 21°C.	21°-24°C.	c. 26°C.

(a) Four samples, laboratory prepared.

(b) Two samples

(c) Ope sample.

Figure in heactet denote huits specified in British Standard Specified in D25 for Cocount Oil; refractive in heactet denote huits specified in British Standard Specified in D25 for Cocount Oil; refractive in heactet denote the property of the Standard Color of the Standard Color of the Palm brief oil; refractive indicated the Standard Color of the St

able from it) depends on the care with which the expression, which contains from 6-10% of all drying process is conducted. In many of the (when prepared in modern mills), and about larger estates outside Caylon hot air drying is 20% of proteins, is used as a cattle food or employed, and in a few places, modern systems fertiliser; the preparation of a press meal or furnace-drying (cotary dners, on sum or turnex-crying (cotary dirers, vacuum dners) have been intailled, which yield a product of very high quality. Strictly aundred copra contains, on the average, 50%, of fat; kuin-dired copra of good quality should contain about 4-6% of water and yield about 50%, of oil with a free fatty and content of less the 10½ (F. R. Combination 1975). of kiln-

over 70% The copra is expressed in oil mills in a manner aimilar to that in which oil seeds generally are worked up. In India, quite good results are obtained in native mills of the pestle and mortar type, but in the Philippine Islands, and other up-to-date tropical establishments, the oil for export is expressed in modern machinery of the same type as is current in Europe or in the United States. Owing to the high oil content, copra is expressed in two stages at a temperature of 60-65°, box-presses [Anglo-American type), cage presses or expellers being employed; the practical yield varies from 63% to 66%, according to the proportion of oil in the copra. The quality of the oil and the proportion of free fatly acids 1 depend upon the

In reporting the free activity of coconut oil, etc.
it is usual to express the percentage of free fatty
activate terms of lauric acid.

(I anchiev, Philippine J. Sci. 1933, 57, 423); tasdiment is not to be confused with any "foot"
or remains of the press-meal, which are normally

austable for human consumption has been proposed (cf. Parker and Brill, Philippine J. Sci. 1917, A, 12, 87).

Coconut oil expressed in the tropies, as well as copra, is now largely shipped in hulk; the copra may be unloaded by mechanical means similar to those employed for discharging grain (/. than 1% (cf. E. B. Copeland, "The Coconut," A. W. Allen, Chem. Met. Eng 1923, 29, 614); 1931). In certain types of kiln-dned or hot air the oil is melted (if necessary) by means of dried copra, the percentage of oil may rise to steam-coils situated in the ships' lanks before

pumping into ahore-tanks,

In temperate chimates, coconut tul is a solid white fat nf bland taste having a peculiar, but ant unpleasant, odour of coconut mest. The odorous aubstances, which are present in very small amount, consist chiefly of methyl heptyl Letone and methyl nonyl ketone (Ifaller and Lassieur, Compt. rend. 1910, 151, 697; Salway, J.C.S. 1917, 111, 407; Brooke, Philippine J.Sci. 1926, 30, 201). These two ketones, together with certain other volatile ketones, carbinols, etc, are removed in the deodorning and refining of cocount oil for edible purposes When crude cocount oil is stored for lorg periods in the liquid condition, a waxy sediment may be formed, from which a purified wax (yielding myricyl alcohol and cerotic acid open quality and condition of the raw material, asponification) can be recovered, which is stated. The press cake obtained as a by-product of the to be austable for use as an ingredient of polishes (Tanchico, Philippine J. Sci. 1935, 57, 423); this tively, by Bömer and Schneider (Z. Unters. by dilute caustic alkalis; on the other hand Nahr. Genussm. 1924, 47, 61), and by Bömer they are so readily converted into soap by and Hüttig (Z. Unters. Lebensm. 1938, means of concentrated caustic solutions that

In consequence of the high contest of sends of low molecular weight (the mean molecular weight of the total fatty acids of coconut oil, for instance, averages 209 as compared with 270-290 for other fats such as palm oil, olive oil, etc.), the yield of glycerol from the saponification of the nut oils (theoretically 13 5-14 1%) is higher than the yield (theoretically 103-11%) from other saap making fats

Refined and deodorsed coconut oil (or palmkernel oil. v. 1nfra) 13 used to an enormous extent as a raw material for the manufacture of margarine, and accounts, in fact, for some 60-66% of the total fats consumed for this purpose in the United States. In Europe, the proportion of coconut, oil employed for mar garina is lower (some 20-30% of the total fata used) owing partly to replacement by palm kernel oil and, especially, to the rapidly increasing use of hydrogenated whale oil

It is also used as an ingredient of shortenings end as a cooking fat ("vegetable hutter"), especially on the Continent, where such preparations as beurre régétal and palmine consist elmost entirely of palm kernel or coconut oil , in the tropics, also, coconut oil is commonly

employed as a cooking oil

Coconul stears a (sodine value 4-7, m p 27°-32°C, asponification value 250-260), which is obtained by filter pressing cooled refined Untern Lebensin. 1925, 55, 354) or rended coconut oil so as to remove the more liquid sumber ("Restrabli"; cf. Grossfeld, ide 193, etc., 194, et glycerides, is used as a chocolate—and con fectionery—fat (qv), Lightly hydrogenated steerin, prepared by pressing inferior grades of the oil, was formerly used in the menufacture of butter fat, which is based upon the microscopical night lights (e CANDLES)

The technical grades of coconut oil are used in enormous quantities in the manufacture of soaps. Owing to the low content of unsaturated acids, and the presence of a large proportion of saturated acids of low molecular weight, the sodium soap of coconut oil (or palm kernel oil, etc) possesses great hardness and the power to retain considerable quantities of water without becoming soft and also a high solohibty and good lathering properties in cold water and in salt water. Further, coconut oil soaps are not thrown out of solution ("salted out") by moderate concentrations of salt, as is the case with the scape of other fats, hence the nut oils constitute the essential raw material for the manufacture of marine scaps for use in senwater. Owing to the hardness, solubility, and ready lathering power of its sosp, coconut oil (or palm kernel oil) is also an important ingredient of most other types of soap, and, especially, of quick-lathering household soaps, flakes, shampoos, shaving soaps, etc., intended for use in luke-warm water.

they are so readily converted into soap by simple mixing of the oil with the equivalent quantity of strong lys at about 40°-50°C. soffices to mitiate the reaction, which proceeds spontaneously to completion with the evolution of heat. Accordingly, coconut oil is extensively employed for the manufacture of soaps by the so-called " cold process "

Minor amounts of coconut oil are used in the manufacture of pharmaceutical and cosmetic preparations, confectionery fillings, and lubri

cants

Owing to its high Reichert-Meissl value, comnot oil, or margarine mixtures containing it. may be used to adulterate butter. Methods for the detection of coconut oil in butter ar mar gamns are based on the detection and estimation of the insoluble volatile fatty acids (caprile and esprie acids) through the analytical deter menations of the Polensie value (cf. Salkowski, anal Chem, 1887, 26, 582; Polenske, Arh Kaiserl, Gesundh amt, 1904, 545; Levio witsch, "Oils, Fats, and Waxes," 6th ed. vol 2, pp 843 et seg); Bertram A and B realues (Bertram, Bos and Verhagen, Chem Weekblad, 1923, 20, 610; Z. deut Oel u Fett Ind 1924, 44, 459, Bertram, ven der Steur, and Verhagen, Chem, Weekhlad, 1925, 22, 849, cf. Kublmann and Grossfeld, Z Unters Nahr. Genussm 1925, 50, 339, (with Baumann), ibid 1926, 51, 27); caprylic acid value (Grossfeld, Z on the hasis of their high content of laune and qualitative test for the detection of coconut oil in appearance of the crystale obtained on re crystallising the fat from ether, is described by Hinks (Analyst, 1907, 32, 160 ; see Lewkswitsch, Ic, pp. 857-858). Similar methods are applied for the detection of coconut oil, etc, in cacao hutter (q v., cf. also Cohn, Z. angew. Chem. 1924, 37, 304; Kulhmann and Grosfeld, 1816, 1926, 39, 24 , Hartel and Marsnus, Z. Unters. Nahr -Genussm. 1924, 47, 205) and soap. Methods for the estimation of butterfut in mixtures con taining coconut oil (e g. margarine, chocolate) depend upon the determination of the content of soluble volatile acids (butyric and caprole acids) in the fat : reference may be made to the methods of Kirschner (Z. Unters. Natr-Genussm. 1905, 9, 65; cf. Bolton, Richmond and Revus, Analyst, 1912, 37, 183; Cocka and Nightingale, shid 1928, 53, 322; Soc. Pub. Analysis, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 64, 405, 84, 415, 1918, 41 Unters. Nahr. Anal. Comm. Rept , sbid. 1936. 61, 408; vin der Laan, Rec. Trav. chim. 1932, 41, 721); of Kuhlmann and Grossfeld (Z. Unters. Lebensm. 1925, 50, 336 : calculation from the Reichert Messel and saponsfication values, cf. I'ntzker, abid 1929, 58, 592); of Kuhlmann and Gross The nut oils further differ from other types of feld (Z Unters Lebensm. 1926, 51, 31; Gross soap oils to that they are not easily apponified feld, abid, 203; abid, 1927, 53, 381; determinaand Verhagen (Lc. supra: determination of Aand B- values).

Since eccount oil and palm-kernel oil are so closely similar, differing chiefly in the amount of insoluble volatile acids present (Polenske value), it is extremely difficult, and may be impossible, to decide by analysis which of the two is present in a mixture of other fats (e.g. margarine), if no information as to origin is available. The determination of the composition of a mixture of the two nut oils is seldom required, as theso fats are rarely adulterated the one with tho other; approximate methods for the purposo depending chiefly upon slight differences in the composition of the "Polenske acids" from the two fats are described by van Kregten, Chem. Weekblad, 1915, 12, 788; Bliehfeldt, J.S.C.I. 1919, 88, 150T; Stokoe, ibid. 1921, 40, 57T; Gilmour, Analyst, 1925, 50, 119; Elsdon, ibid.

1927, 52, 63.

Palm-kernel Oll is obtained from seed-kernels of the oil-palm Elacis guinecusis Jacq., which is indigenous to West Africa but also cultivated in the Dutch East Indies, Malaya, and other tropical countries. More than 95% of the palm kernels entering world-trade, however, come from Africa—British West Africa, French W. and Equatorial Africa, and tho Belgian Congo being the principal producing areas; in the Asiatic regions the palm kernel industry is subordinate to the production of palm oil, and of minor importance at present. The "palm nuts" (i.e. the "stones" of the drupaceous fruits) are separated from the oleaginous fruit-pulp when the latter is worked up for the production of palm oil. Tho "nuts" are dried by exposure to the sun or in kilns sufficiently to loosen the kernels (seeds) within the shells, and the comparatively thin shells are cracked by nativo hand-labour or by machinery. The kernels are then separated from the shell-fragments by screening or by flotation in brine or clay-water (sp.gr. 1-1-1-5), dried if necessary, and exported in bulk to Europe or the United States. Unlike palm oil and ecconut oil, palm kernel oil is not prepared (except for local or native consumption) in the palm-growing regions, the exports being confined to the shelled kernels. In Europe, etc., the kernels are erushed and expressed in the usual way in two stages at 45°-60°C., yielding 43-45% of oil on the largo scale. A certain amount of oil is also recovered (in Europe) by extraction with solvents.

The press-cake contains 6-8% of oil and about 2.5% of nitrogen: it is employed in the feeding of dairy cows and other farm animals (cf. "Use of Oil Cakes and Extracted Meals," Ministry of Agriculture and Fisheries, Bull. No. 11, 2nd ed., 1937; Anon., Bull. Imp. Inst. 1915, 13, 151, 446; C. Crowther et al. J. Agric. Sci. 1917, 8, 429, 451; W. Godden, ibid. 419).

Palm-kernel oil is white to pale yellow in colour and resembles ecconut oil in composition (cf. Tables I, II, III), appearance, and odour; as in the case of coconut oil, the characteristic aroma appears to be due largely to the presence of methyl nonyl ketone (Salway, J.C.S. 1917. of methyl nonyl ketone (Salway, J.C.S. 1914, 111, 407). It is used for the same purposes as they were frequently wrongly attributed to the fibre coconut oil; pressed "palm nut stearin" palm ("plassava") Attalea funifera Mart.

tion of butyric acid value): and of Bertram, Bos like the corresponding "coconut stearin." is employed as a chocolate fat, and the "palm nut olein" for soap making. Palm kernel oil is generally regarded as slightly inferior to coconut oil and, as a rule, commands a lower price; the choice between the two fats is, however, largely influenced by geographical and politico-economie considerations and trading custom. Thus, before the Great War, Germany held a virtual monopoly of the palm kernel erushing industry, exporting the surplus oil to the United Kingdom and other countries, and since 1925-26 has recaptured her position as premier producer and consumer of the oil, in spite of the growth of important palm kernel crusbing industries in the United Kingdom, Franco and other countries. Imperts of palm kernels have been of the order of 250,000-300,000 tons per annum from 1927-1935 (374,000 tons in 1936 and 322,500 tons in 1937) into Germany, and 130,000-160,000 tons per annum into England over the same period (c. 200,000-250,000 tons per annum from 1920-1926); French imperts of the kernels, which were only of the order of 12,000 tons per annum in 1928-1933, have latterly shown a notable increase to 119,400 tens in 1936 (69,000 tens in 1937). Total imports of kernels into the principal importing countries have been 594,200, 735,800, and 675,050 tons in 1935, 1936, and 1937 respectively (cf. "Review of Oil and Fat Markots," 1937, H. M. Fauro & Co.; for a detailed survey up to 1932 of the production and trade in palm kernels and palm kernel oil in British and fereign countries, of. "Survey of Vegetable Oil Seeds and Oils, vol. 1, Oil Palm Products," Empire Marketing Board, E.M.B. 54, 1932, H.M.S.O.).

For the determination of palm kernel oil in mixtures, see under Coconut Oil.

Babassu Oil is obtained from the kernels of the Brazilian babassu palm, Orbignya speciosa Barb. Rodr. (=Orbignya Martiana),1 Tho woody endocarp ("shell") is extremely hard, and considerable difficulty has been experienced in devising machinery capable of splitting this shell without eausing excessive damage to the 2-6 small kernels (which only weigh about 4-6 g. each) contained therein. The kernels contain from 63-70% of fat (cf. Tables I, II, III), which closely resembles eoconut oil and is used for similar purposes. The woody shells may be used as a fuel, or subjected to destructive distillation for the recovery of acetic acid, etc., and metallurgical charcoal. Babassu fat is one of the principal native fats of Brazil, and since about 1915 considerable quantities of the kernels have been exported to Europe, and, more recently, to the United States. As in the case of the coconut, the press-cake is suitable for use as cattle-food (cf. Anon., Bull. Imp. Inst. 1929, 27, 286; "The Babassu," Ministry of Agric., Rio de Janeiro, 1930; Arié, Arch. Inst. Biolog. São Paulo, 1931, (4), 301; Heiduschka and Agsten, I.c., Table II).

Cohune Nut Oil, from the seeds of Orbignya cohune Mart. Dahlgren (=Attalea cohune

Mart.) indigenous to British Honduras and Central America, has so far been of little commercial importance, owing largely to lack of development on the agricultural side of the industry, and to the difficulties experenced in providing machinery in place of laborious hand-labour to crack the small hard nuts. The services amountage 42, 32, 50% of the blook services amountage 43, 50% of the blook services are supported by the services of the latest providing the services of the services of the high services of the service

Murumuru Fat, from the thin-shelled seeds of another Brazilian palm, Astrocaryum murumuru Mart., is a haed, hrittle fat resembling coconut stearm, and is similarly employed as a

ehocolate fat.

Although it may be included among the not oils for convenence, it will be seen from Table II that murumura fat contains a much smaller proportion of caprylic and eappre acids then do the typical fats of the group (the Reichert-Messe), and not have a supported by the second of the theory of the second of the s

coryum tuciuma Mart. (Collin, Biochem, J. 1933, 27, 1369), and Aerce catcha' (the areca or betelnut paim) (Rathje, Areb. Pharm 1908, 246, 702) have a high content of laune and myrastic acids, but are comparatively poor in the lower acids; these fats have no commercial significance other than for local consumption in the

countries of origin.

Khakan Fat, which is chemically very similar to coconut oil, is not derived from a paim but from the unrelated plant Salvadora ofeoides Done (fam. Salvadoraceæ), and will be discussed under Khakan Fat.

COCOSITOL, mp. 342°, optically inactive, a natural mactive form of inositol, is identical with scylitol and queraine (Riugo Müller, J.C.S. 1912, 101, 2383). It was found by him in the leaves of two species of Cocos (C. plumoza and C. nucifero). The term scylitol only is retained.

COOAMINE, C2041, O,N, m. p. 128-127. (bexagonal prasms from £1,0), is a minor open alkaloid. The base was first noisted from opum by llesse (Annalen, 1870, 135, 66). It reacts strongly alkaline, is moderately soluble in Holl. It gives a grand very soluble in E1041. It gives a gree colour with ferric chloride, and nitrie and dissolves the alkaloid forming a dark green byud.

Spath and Epstein (Ber. 1926, 59, [B], 2791) obtained d-laudanosine by methylation of codamine and have conclusively demonstrated that the alkaloid is a laudanosine derivative with a phenole OH-group in 7-position

CODEINE, C₁₈H₁₁O₃N,H₁O, an isoquinolme alkaloid, exists in Turkey opium in the proportion of about 0.75%, while in the Persian or Indian varieties the average is about 2.25%. COOEONAL. Combination of codeme

COGEONAL. Combination of codeme dethylbarhiturate, 2 parts, and sodium diethylbarbiturate, 15 parts, in tablet form. It is a sedative and hypnotic (Knoll, Ludwigehafen, Pharmaceutical Praducts, London). B.P.G.

COO-LIVER OIL, in the narrowest sense, is the oil obtained from the livers of the European and American cod-fish, Gadus morrhus L. (G. caillaras), and the Japaneses cod (" madara " or " tara"), G. macrocephalus Tilesus; in

or "ara"), G. macrocephalus (Riemus; mordinary prastice, however, which is anotioned by the definitions of the pharmacopous of the warnous countres, the term "cod liver oil" includes the equivalent liver oils from allied species of Gadus or of the family Gaddae, e.g. the pollack (G. pollachius), the coal fish or author, hake (Merinecius medicardae), ling (Moltamoto), the Spaneer Pollachius Brandin, Jordan mosto), the Spaneer Pollachius Brandin, Jordan mosto), the Spaneer Pollachius Brandin, Jordan challengeman Pallas (Gadus chalcopramma)—se caught in Isrgo quantities, but up to the present the here-oil has not been used for medicanal purposes.

The coal appear annually at certain seasona

of the year in enormous shoals in the constal regions of the northern sess in latitudes shove 45°N.; and, correspondingly, important fishing grounds have grown up on the coasts of Norway, Scotland, Iceland, Newfoundland, on both eastern and western coasts of Canada and the United States, and off the shores of Japan and Siberia, As a result of the introduction of factory ships, even more remote fishing-grounds, such as those at Bear Island, Spitabergen, can be developed for the production of first-class oils. Whilst the oldest and probably the bestknown commercial fishing grounds are those of Norway, whence cod liver oil (" Bergentran ") was exported (for leather dressing) as early as the fifteenth century, the Newfoundland industry is also of considerable antiquity, and records exist of the export of "train oil" (codoil) to England from the seventeenth century. This area is now becoming increasingly important for the production of high grade medicinal oils, the bulk of which are exported to Canada and the United States. Both in Newfoundland and Norway, the preparation of medicinal cod hver oil is conducted under strict Government supervision. The production of medicinal oils on the east coast of the United States and in Japan is also considerable. Until recently the interests of the English native industry were restricted to the technical " cod " oils, but of late years efforts have been made to promote manufacture of domestic medicinal ods.

In the Newfoundland and Lofoten fisheries of
Norway the catch consists exclusively of eod,
but in the Finmark fishery of Norway and
the English coastal fishery, sauthe, pollack,
bake, ling, and other closely ellied members of
the cod family are included. The liver oils of
Schl. these fish, however, are closely similar in

ehemical composition to that of the true cod, I and since they are equal to, or in some cases superior to, pure cod-liver oil in vitamin potency, a well-nourished state.

In the oldest process for the recovery of eodliver oil, the livers, after removal from the fish, were heaped up in barrels and allowed to undergo spontaneous putrefaction in order to decompose the liver cells and permit the oil to exude, which was drawn off at intervals as the rotting proceeded, in fractions ranging from pale brown to almost hlack in colour; the process might he accelerated hy exposing the barrels to the sun. Some improvement in the quality of the oil was effected hy careful selection of sound livers, and the removal of blood of the gall-hladders and other visceral matter before treatment, hut, nevertheless, even the best oils prepared by the "rotting" method were contaminated with decomposition products of the liver tissue (organic hases) and possessed a rank, fishy odour and nauseous taste.1 This crude method of preparing the oil is now practically obsolete and, except in a few primitive installations producing low-grade oils, has been replaced by the "steaming process." In the original process introduced by Charles Fox in Newfoundland (1848, cf. Munn, Montreal Pharm. J. 1914, 25, 192) and developed hy P. Möller in Norway, 1853 (cf. Möller, "Cod-liver Oil and Chemistry," 1895), the fresh, sound, washed livers were rendered in steamjacketed pans, hnt to-day it is more usual to "cook" the livers with open steam (at a pressure of from 30-100 lb./sq. in.) which is blown directly into the mass of fresh livers contained in an open or covered pan.

For the production of the finest (medicinal) grade of oil, tin-lined pans or vessels of noncorroding metal, are employed, and only sound livers in the freshest possible condition are taken; all diseased or discoloured livers are rejected and worked np separately for the technical grades of oil. The gall-hladders are removed, and the livers are thoroughly washed hefore steaming, which is continued for a sufficient time (ahout 30 minutes) to ensure the destruction of any fat-splitting enzymes, which might otherwise cause deterioration of the oil during subsequent storage. The oil which separates freely from the livers is dipped off, strained, and allowed to stand in order that traces of liver tissue and moisture may settle out, and the purification is completed, if necessary, by filtration, centrifuging, or treatment with fuller's earth; finally, the oil is packed into tin-lined barrels for transport or storage. In order to prevent deterioration of the oil and loss of vitamin-potency through oxida-

1 Details of the "rotting" process and the early steaming process are described by Möller, "Cod-liver Oil and Chemistry," London, 1895; cf. also Lewkowitsch, "Oils, Fats, and Waxes," 6th ed., vol. II,

tion, exposure of the oil to air and light is minimised throughout the operations. further details of the preparation of medicinal superior to, pure cod-liver oil in vitamin potency, no objection to their presence is made. In the Lofoten fishery the fish are taken during the spawning season in a fasting condition, and the livers are poorer in oil than those ohtained from the fish which are canght in the Finmark or "Lodde"-fishing season, and off Newfoundland, when the cod is feeding actively and in land, when the cod is feeding actively and in Rev. Trav. Off. Peches marit 1930, 3, 1030; and land properties of the preparation of medicinal cod-liver oil in Norway, Newfoundland and Iceland, cf. Zilva and Drummond, J.S.C.I. 1922, 41, 280T; 1923, 42, 185T, 250T; Drummond and Hilditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Hilditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values of Cod-liver oil in Norway, Newfoundland and Holditch, "Relative Values o Chahre, Thesis, Paris, 1936; Macpherson, Bull. 3, Fish Sect. Dept. Nat. Resources, Newfoundland, 1937).

The medicinal oil obtained in this manner is consumed as " congealing oil " in those countries where the pharmacopæia or popular taste raises no objections to such an oil, which deposits a solid "stearin" when exposed to low temperatures. In England and certain other countries, however, the pharmaceutical trade prefers, and the pharmacopæia prescribes, a "non-freezing" oil, which does not set at low temperatures; this is prepared by "refining" or chilling the crude oil at temperatures between 0°C. and -10°C. in order to precipitate the higher-melting glycerides ("stearin") which are filtered off at about 0°C., leaving a yield of about 75-80% of a clear "non-freezing" oil. The separated "stearin" (consisting actually of mixed glycerides of saturated and unsaturated acids) may be pressed in order to recover any entrained liquid oil (which is added to the second grade of oil) and is sold as the best quality "fish stearin ' for soap manufacture.

The steamed liver-pulp remaining in the rendering pan may be holled up again with steam before it is finally pressed, yielding in both cases oils of good quality which are employed as a vitamin-active supplement in the feeding of poultry and cattle. The final residue of liver-tissue is dried and converted into livermeal for animal feeding. In general commerce, the expression "cod-liver oil" is restricted to the medicinal oils intended for human or animal nutrition, whilst the term "cod-oil" ("hrown eod-oil," "hrown oil") is used for the technical oils which are obtained by steam-rendering (or, more rarely, by the rotting process) from livers which, on the score of condition or freshness, are unfit for the production of medicinal oil; where other fish besides cod are included in the catch, the oil is known commercially as "coast cod oil." The cod-oils, in general, have a high acid value and rank odour, and are contaminated with impurities derived from the decayed livercells; the use of such oils for the feeding of animals is to be deprecated, especially as there is some evidence that oils from putrid livers may he definitely toxic. The crude "nnracked" cod-oils contain considerable cod-oils contain considerable quantities of stearin which are removed hy chilling and filtering or settling, and used in soap-making or as "fish-tallow" in leather Cod-oils find extensive use in the currying. leather industries, where the raw oil is employed in the chamoising process and in the currying (stuffing) and oiling of heavy leathers; further, considerable quantities of sulphonated (sulphated) cod-oil, e.g. Turkey-red oils, which is prepared by treating the oil with concentrated

aulphuric acid are employed for the oiling-off deeper yellow colour than the Norwegian oils. of heavy leathers and for the fat-luquoring of [N.B. This inherent colour is quite distinct light leathers and chrome-tanned leathers, from the discoloration of oils which have Technical cod oils are also employed in the been prepared from stale hvers or by had manufacture of lubricating greases and low-methods.) grade soaps. The dried, pressed liver meal ture is utilised as "fish manure."

uality are pale in colour and have only a abghtly livers from fish of both sexes, different size, fishy odour and tastes; in consequence, probably, of the different feeding conditions, the olar from flowfould livers of fails of different size, Newfoundland and Iceland possess a distinctly etc., show wider extremes of variation.

The analytical characters of medicinal codrecovered as a hy-product of cod oil manufac- liver oils are illustrated in Table I, except where otherwise stated, the figures refer to oils pro-Pure medicinal cod-liver oils of pharmacopmia duced on the large scale from hatches of mixed

TABLE I. CHARACTERISTICS OF CONTAVER OUR

	Medicinal olla (mostly non congealing),1	Pharmacopæisi requirements (non congesting oils), ^a	Crude whole (s.e. congeal- lng) medi- cinal olla	Pure cod liver oils	Japanese cod- liver (madara) plis s
Sp gr	0 920-0 9289/ 25°C.	0 922-0 929/ 15 5°C, (B.P.) 0 918-0 927/ 25°C. (U.S P.)		0 924-0 929/ 15 5°C.	0 9243-0 9397/ 15°C. (0 924-0 9298)
Ref. index	1·474–1·4798/ 20°C.	1-4705-1-4745/ 40°C, (B.P.)	_	1 4705- 1 4728/40°C.	1 4742-1 4803/ 20°C (1 4765-1 4808
Saponification value	175–197 (usually 180–190)	180-190 (B.P.) 180-192(U.S.P.)		181-189	174 5-190 7 (183-189)
Iodine value ,	147-183 (usually 150-170)	155-173 (B.P.) 145-180 (U.S.P.) 4	153-162 (a) 142-185 (b) 159-174 (c) 140-185 (d)	154-172	143-205 (152-182)
Acid value .	05-35	⇒1.2 (B.P.) ⇒28 (U.S.P.)	06-24	_	0 4-5 0 (0 5-12)
Uneaponifiable matter . ,	0 9–1-7%	1.5% (B.P.) 1.3% (U.S.P.)	- /	0 96-1-52%	0 8-4-5% (0 6-1-0)

1 (f Lewkowitsch, "Olis, Fata, and Wases," vol. II; A. D. Holmes, Ind. Eng. Chem. 1924, 16, 1181; Indians and Cloud, Oil and fat Ind. 1927, 4, 603; Ender and Jermsted, Harm Zentrah. 1907, 77, 104. Indians and Cloud, Oil and fat Ind. 1927, 4, 603; Ender and Jermsted, Harm Zentrah. 1908, oil her oil is from an Oil from Jermster, oil her oil is from an Oil from Jermster. Oil her oil is from an Oil from Jermster. Oil her oil is from an Oil from Jermster. Oil her oil is from an Oil fato, Indians, John St. Jermster. Oil from Jermster. Oil from an Oil fatol, Indians, Charles, Cha

expected, somewhat higher lodine values and pharmacopocial oils, but usually possesses a refractive midnes than the crude oils from stronger findly taste and alightly darker colour, which they have been derived (cf. Holmes and Pigott, Ic.; Kwazi, Ic.; 1933), but the difference of the within the range of variation abown by Drummond and Hidtich, Ic.; for review of the terude oils. There is no significant different oil leteratures of Hidtich, Ic.; for review of the increase of the result of

The non-freezing oils have, as would be consumption is only slightly inferior to the expected, somewhat higher lodine values and pharmacoposial oils, but usually possesses a cod oils have lower sodine values, depending upon The acid value of the best oils is frequently less the amount of oxidation the oil has auffered, than 05; the second quality of oil for animal and high free aciditics. The following figures (due to Holmes, Clough and Owen, Oil and Fat | higher percentages of unsaponifiable matter Ind. 1929, 6, (No. 10), 15) for twenty oils of may be found in coast cod oils. various grades, including sun-rotted oils, are illustrative: sp.gr. 0.9193-0.9266 at 25°, $n_{\rm D}^{20}$ 1.2-2.8%, free fatty acids 7-20.2%. Somewhat included for comparison:

The approximate quantitative composition

of the fatty acids of cod-liver oil is indicated in 1.480–1.483, saponification value 184–194.5, Table II, in which the corresponding figures for iodine value 146.3-168.6, unsaponifiable matter some allied and closely similar-fisb-liver oils are

TABLE II. PERCENTAGE COMPOSITION OF FATTY ACIDS OF FISH LIVER OILS (Guha, Hilditch and Lovern, Biochem. J. 1930, 24, 266; cf. Drummond and Hilditch, l.c.)

Fatty acids.		Liver oils.		
No. of carbon atoms per mol.	Cod.	Saithe.	Hake.	Ling.
Saturated Acids. C ₁₄	$ \begin{array}{c cccc} (a)^1 & & & (b)^2 \\ \% & & \% & & \% \\ 3 \cdot 5 - 6 & & 2 & \\ 6 \cdot 5 - 10 & & 14 & \\ 0 - 0 \cdot 5 & & 1 & \\ \end{array} $	% 6·5 12–13 0–0·5	% 4·5-7 11-13 0·5	% 5 13 1
Unsaturated Acids. C ₁₄ (—2H) C ₁₆ (—2H) C ₁₈ (—3H) C ₂₀ (—xH) C ₂₂ (—xH)	$ \begin{array}{c cccc} 0-0.5 & & & 2 \\ 16-20 & & & 10 \\ 25-31 & & 26 \\ 25-31.5 & & 25 \\ (x=5-6) & & 25 \\ 10-14 & & 20 \\ (x=7) & & <1 \\ ? & & <1 \\ & & (x=6-6) \\ & & & (x=7) \\ & & & <1 \\ \end{array} $	$\begin{array}{c c} (x=5) & (x=5) \\ 10-16\cdot 5 & (x=7) \\ ? & ? \end{array}$	trace 8-17 18-34 25-31 (x=4-5) 14-16 (x=6)	trace 13 32·5 24 (x=6) 11·5 (x=7) ?

Figures in brackets denote average unsaturation of each fraction, e.g. (-2H) denotes one double bond; (-3H), a mixture of perhaps mono- and di-ethenoid acids, averaging one and a half double bonds.

Range of analyses of Scotch, Norwegian, and Newfoundland cod-liver oils.

North Sea cod-liver oil: average unsaturation of the C₁₈ unsaturated acids = -3·3 H. Harper and Hilditch,

J.S.C.I. 1937, 56, 322T.

The saturated fatty acids (10-15% of the total | lower saturated (volatile) fatty acids, such as acids) consist chiefly of myristic and palmitic acids, with traces of stearic acid (Guha, Hilditch and Lovern, I.c.; Toyama, J. Soc. Chem. Ind. Japan, 1926, 29, 172B). The unsaturated acids Japan, 1920, 29, 1725). The insaturated acids include a large proportion of the monoethylenic zoomaric (Δ^{9.10}C₁₆H₃₀O₂), oleic (Δ^{9.10}C₁₈H₃₄O₂), gadoleic (Δ^{9.10}C₂₉H₃₈O₂), and cetoleic (Δ^{11.12}C₂₂H₄₂O₂) acids (H. Bull, Ber. 1906, 39, 3570; Toyama, J. Soc. Chem. Ind. Japan, 1926, 29, 624; 1927, 30, 154B, 155B; 1934, 37, 148b; projected: 37, 14B); myristoleic ($C_{14}H_{26}O_2$), and selacholeic ($\Delta^{15.16}C_{24}H_{46}O_2$) acids are present in minor amount (Toyama and Tsucbiya, ibid. 1935, 38, 684B). More highly unsaturated acids of the C₂₀ and C₂₂ series, including docosabexaenoic acid (C₂₂H₃₂O₂) (cf. Farmer and Van den Heuvel, J.C.S. 1938, 427), are also important constituents; stearidonic acid (C₁₈H₂₈O₂) and arachidonic acid (C₂₀H₃₂O₂) bave been identified by Suzuki (Proc. Imp. Acad. Tokyo, 1929, 5, 165) and nisinic acid (C₂₄H₃₆O₂) by Toyama and Tsuchiya (J. Soc. Chem. Ind. Japan, 1934, 37, 531B). The "jecoleic acid" and "therapic acid" of the early investigators (cf. Heycrdabl, in F. P. Möller's "Cod Liver Oil and Chemistry," 1895) were probably more or less impure preparations of gadoleic and stearidonic acids respectively.

acetic, butyric, and capric acids, as well as of organic bases in cod-liver oil were caused by the presence of decomposition products of the oil and liver tissue in the old "rotted" oils.

The unsaponifiable matter of cod-liver and cod-oils amounts normally to 0.7-2.8%,1 although a much larger proportion has been reported in individual cases of the oils from single lean livers of old, large fish (cf. K. Kawai, J. Pharm. Soc. Japan, 1933, 53, No. 10, 183); as a rule, about half the unsaponifiable matter; consists of cholesterol, the remainder comprising the fat-soluble vitamins (see below), colouring matters, and other non-sterol constituents including squalene and batyl alcohol (Drummond, Channon, and Coward, Biochem. J. 1925, 19, 1047), one or two other unsaturated hydrocarbons, and, perhaps, other mono- and di-hydric (aliphatic) alcohols (Wicdemann, ibid. 1932, 26, 264; Kawakami, Bull. Inst. Phys. Chem. Res. Tokyo, 1933, 12, 825, 834, 845 (Abstr. 7, 8, 9, publ. with Sci. Papers I. P.-C. R. Tokyo, 1933, 22, No. 457-467).

Cod-liver oil, even of unexceptionable quality, contains a small amount of iodine, of the order

ad Chemistry," 1895) were probably more or simpure preparations of gadoleic and stearionic acids respectively.

The older reports concerning the occurrence of the unsaponifiable matter (cf. S.P.A. Sub. Comm. Rept., Analyst, 1933, 58, 203).

of 3-15 pts /million (cf. Hopfer de l'Orme, | thereof, such as carotene), which is present to the Hufeland's J. 1836 (April), p. 115; Heyerdahl, extent of some 100-500 pts /million, chiefly in lc.; Holmes and Remington, Amer. J. Ds. estrified form (Bacharach and Smith, Quart, J. Chem. 1934, 26, 573) and Luzanski (Tids Kiemi, 1936, 16, 56).

In common with other fish-liver oils, cod liver oil gives a red violet coloration when concentrated sulphure acid is added to a solution of the oil in chloroform or carbon disulphide, the depth of the colour varies with different oils, and appears to be due to a condensation reaction of some micor constituent of the oil (hpochrome *) associated with vitamin-A and analogous to the antimony trablonde colour reaction of the latter (c. 1sfra) (cf. Drummond and Watson, Analyst, 1922, 47, 341). With Japanese and clay, a blue green coloration as developed (cf. Kobayashi, Mem. Fac Sci. Eng. Tolyo, 1927, 4, 23). These colour reactions may fail, however, with hydrogenated (hardened) liver oils.

A green colour reaction is observed when a solution of cod liver (or other fish liver) oil in glacial acetic acid is treated with a mixture of hromine and chloroform and sodine bromide (Hanus iodine solution) (Tortelli-Jaffe reaction . Tortell and Jaffe, Chem Ztg 1915, 39, 14, cf. Better and Szimkin, Fettchem. Umschau,

1934, 41, 225). Although cod liver oil seems to have been held in esteem as a domestic remedy for many centuries among the inhabitants of the northern coasts, its systematic use in the treatment of bone disorders, tuberculosis, and conditions of malnutration appears to date from the successful clinical trials made at Manchester in the second half of the eighteenth century by Dr. Kay and his colleagues (cf. T. Pereival, Medical Essays, 1789, 11, 354), and despite the nauseous character of the oils available at that time, the popularity of cod liver oil as a remedy steadily increased through the nineteenth century. The extensive modern use of cod liver oil in the prophylaxis and cure of ricketa and other accepted) conditions associated with defective calcium. The premetabolism is now so well known as to be a by the blue colour (absorption maxima at λ = commonplace. The remarkable therapeutic 572-584 and 617 mm, which is obtained on properties of the oil, which have been variously treating a solution of the oil in chloroform ascribed to the presence of organic bases, to with antimony trichloride (Carr-Price reagent, the presence of traces of sodine, and to the Biochem J. 1926, 20, 497; cf Analyst, 1931, 56, the presence of traces of sodine, and to the Brochem J. 1926, 20, 497; cf Analyst, 1931, 56, specific nature of the highly unsaturated acids 457; 1932, 57, 302; "British Pharmacopoua," which compose the hull, of the glycendes of the 1932). The intensity of the roloration (which may oil, are now attributed primarily to the presence, in comparatively high concentration, of the fat-

Children, 1935, 49, 94). The presence of traces Pharm. 1928, 1, 539; Reti, Compt. rend. Soc. of arsence (from 1-4 to 5 1 pts./million) in unbiol. 1935, 120, 577; Hickman, Nature, 1936, refined cod liver oil from fresh livera has been 133, 881; Ind. Eng. Chem. 1937, 29, 1107; reported by Holmes and Remington (Ind. Eng. Hamano, Sci. Papers Inst. Phys. Chem. Res. Tokyo, 1937, 32, 44). Its presence is indicated by a maximum at $\lambda=328$ m μ in the absorption spectrum of the oil or of the unsaponinable fraction therefrom (K. Takahashi, et al. J. Chem. Soc. Japan, 1922, 43, 826; 1923, 44, 590; Sci. Papers Inst. Phys. Chem Res. Tokyo, 1925, 3, 81, Heilhron et al Biochem. J. 1928, 22, 987; 1931, 25, 20; J.S.C 1. 1931, 50, 183T), the intensity of which affords an approximately the intensity of which affords an approximately quantitative measure of the vitamin-content of the oil (cf. Coward, Dyer and Morion, Biochem. J. 1932, 26, 1993; Carr and Jewell, Nature, 1933, 132, 92; Gillam, Heilbron, Morton, Bishop and Drummond, Biochem J. 1933, 27, 878, Booth, Kon, Dann and Moort, deed, 1189; Baumann and Steenbock, J. Boil Chem. 1933, 101, 547; Chevaller and Chahre, Biochem J 1933, 27, 298, Jones and Christian-sen, J. Amer Pharm. Assoc 1935, 24, 1072). This test (the examination being conducted on a solution of the unsaponifiable matter) was recommended as a method of vitamin A assay by the Second International Conference on Vitamin Standardisation (1934), and has been admitted as a subsidiary test in the 1936 Addendum (Appendix IV) to the "British Pharmacopeaia" of 1932, where it is regarded as giving a trustworthy indication of the vitamin-A content of cod liver oils which conform in all other respects to the requirements of the "British Pharmacopeia" (As there may be other substances, however, which show a aimilar absorption in the region of \$\alpha = 328 m\mu, the test may be inapplicable to certain vitamin preparation, etc., and the "British Pharma-copena" Addendum prescribes that, in the event of any divergency between the results of the vitamin A assay by the spectrophotometric method and by the hological method, the value as determined by the latter method should be

The presence of vitamin-A is also revealed be measured spectrophotometrically, or by meana of a colorimeter under standard conditions) in comparatively high concentration, of the fat-soluble viriammes-I and .D (cf. Obborns and smarr soughly parallel to the viriamm activity of Mendel, J. Biol. Chem. 1914, 17. 401; Drum-bond, Biochem. J. 1910, 13, 81). (It may be oil as measured by hiological methods, attracted renewal attention as a source of inhibitors in the oil, or perhaps to the presence combined todine in the treatment of certain of an allied but hlotography anactive chromogen deficiency conditions, cf. Chadester and others, (cf. O. Rych, Bull. Soc. Chim. biol., 1921, 188, Nature, 1932, 130, 541; Science, 1932, 76, 609; Ender, Biochem. J. 1932, 151, 153, Lovern and others, ideal, 1933, 51, 181, 182. The vitamin-A activity of cod liver oil results Better correlation is obtained if the colour from the occurrence in the oil of the actual test is performed on a solution of the unsaponivitamin-A itself (as distinct from a precursor fiable fraction from the oil (cf. Smith and

Analyst, 1932, 57, 368; Emmerie, Nature, 1933, 131, 364; 1935, 136, 183). (It may be noted, however, that this colour test, which figured in the 1932 edition of the "British Pharmacopæia" has been discarded in the 1936 Addendum in favour of the spectrophotometric test.) For further details concerning these tests, and of the hiological assay of vitamin-A, see "British Pharmacopæia" Addendum, 1936. An interesting modification of the Carr-Price test, in which a less fugitive violet colour (λ_{max} =545 and 475 m μ) is obtained by the use of guaiacol as reagent, is described by E. Rosenthal and Erdélyi (Biochem. Z. 1933, 267, 119; 1934, 271, 414; Biochem. J. 1935, 29, 2112; Rosenthal and Weltner, *ibid*. 1935, 29, 1036).

The vitamin-A activity of cod-liver oil varies within wide limits according to the age and sexual condition of the fish, its place of origin, and its food; eases of oils containing from 210 to 3,400 International Units (I.U.) per gram arc reported by Chevallier (Ann. Chim. Analyt. 1936, (iii), 18, 93; cf. Chabre, Thesis, Paris, 1936; cf. K.Kawai, J. Pharm. Soe, Japan, 1932, 52, No. 10, 95; No. 12, 169); the "British Pharmacopæia" Addendum (1936) and the "United States Pharmacopæia," XI (1936), impose a minimal requirement of 600 I.U. per g. of oil. The figures of Chevallier and Chabre, like those of Drummond and Hilditch (E.M.B., No. 35, l.c.) support the view that, on the whole, the oils from Newfoundland and Iceland are richer in vitamin-A than the Norwegian oils. On the other hand, the Norwegian (Lofoten) oils appear to be richer in vitamin-D (cf. Poulsson, Nord. Med. Tidsk. 1933, 6, 886; Amer. J. Pharm. 1934, 106, 102). The vitamin-D content of codliver oil also varies considerably, and is quite independent of the amount of vitamin-A present (cf. Coward, Dyer and Morgan, Analyst, 1932, 57, 368; van Esveld, Nederl. Tijds. Geneesk. 1935, 79, 2924; Kawai, J. Pharm. Soc. Japan, 1934, 54, No. 5, 87). On the whole, the less the oil content of the cod liver; the greater is the concentration of vitamin-D in other words, the depletion of the fat reserves in the liver during spawning, fasting, or ageing of the cod-fish is not accompanied by a corresponding drain on the store of vitamin-D (cf. Poulsson, l.c.; Kawai, l.c., 1934).
According to K. Kawai (l.c.) the same tendency

is observable in the case of vitamin-A and the blue (Carr-Price) ehromogen, very bigh potencies, and especially high blue values, being observed in the case of oils from livers which, owing to the age or condition of the cod, were very poor in

Before the anti-racbitic principle (vitamin-D) of eod-liver oil had been isolated it became clear from biological comparison of the relative therapeutic efficiencies in tests with different animals (rats and chicks) that the active agent in cod-liver oil was not identical with calciferol (vitamin-D₂), obtained by the irradiation of ergosterol (cf. Massengale and Nussmeier,

Hazley, Biochem. J. 1930, 24, 1942; Church and Wileox, ibid. 1934, 107, 735; J. Nutrition, and Norris, J. Biol. Chem. 1930, 87, 139; 1935, 9, 569; Bills, J. Biol. Chem. 1935, 108, Coward and others, Biochem. J. 1931, 25, 1102; 323); and it has recently been shown by Zueker, Simons, Colman and Demarcst (Naturwiss. 1938, 26, 11) that the vitamin-D of cod-liver oil, like the vitamin-D which has been isolated (via the dinitrohenzoate) from tunnyliver and halihut-liver oils (Brockmann, physiol. Chem. 1936, 241, 104; 1937, 245, 96; cf. Simons and Zucker, J. Amer. Chem. Soc. 1936, 58, 2655), is in all probability identical with "vilamin- D_3 " prepared artificially by the irradiation of 7-dehydrocholesterol (Windaus, Schenk, and Werder, Z. physiol. Chem. 1936, 241, 100; Grah., ibid. 1936, 243, 63; Folherth, Linderheilk. 1937, 59, 329). Da has been obtained in the crystalline state (Schenck, Naturwiss. 1937, 25, 159) and it bas heen shown that, despite the difference in the ehemical constitution (the side-chain of the molecule), vitamin- D_3 has the same antirachitie potency (40 million I.U. per g.) as ealciferol when tested on rats (International method). Like the rat, the human subject appears to be equally sensitive to both forms of vitamin-D, that is to say, the administration of an equal number of units (as measured by the usual tests with rats) of calciferol, vitamin-D₃, or codliver (or other active fish-liver) oil produces quantitatively the same therapeutic effect (cf Eliot, Nelson, Barnes, Browne and Jenss, J. Pediatrics, 1936, 9, 355, 761). The chick, on the other hand, discriminates between the different vitamins, for, whilst it readily utilizes vitamin-D₃ or the cod-liver oil vitamin, it is most inefficient in its use of calciferol, so that some fifty times as many (rat-)units (I.U.) of calciferol are needed to evoke the same response as any given number of units of cod-liver oil or vitamin- D_3 . The possibility that more than one form of vitamin-D may occur in fishliver oils is mooted by Bills, et al. J. Amer. Med. Assoc. 1937, 108, 13; J. Nutrition, 1937, 13, 435).

> The vitamin-D of cod-liver oil appears to be present partly in the free state and partly as esters (Hickman, Nature, 1936, 138, 881; Ind. Eng. Chem. 1937, 29, 1107). The amount in the oil is probably of the order of 2-8 parts per million, but up to the present only biological methods of comparing the potencies of different oils have been adopted (cf. suggested method in "British Pharmacopæia" Addendum, 1936, Appendix XV, also article VITAMIN-D); hoth the "British Pharmacopœia" Addendum and the "United States Pharmacopæia" (1936) require a minimum potency of 85 I.U. per g. of oil. A photometric method for the assay of vitamin-D, which has been proposed by L. Fuchs and Beck (Pharm. Presse, Wiss.-Prakt. Heft, 1933, 38, 81, 93) requires further investigation.

The origin of the vitamins in cod-liver oilwhether they are merely ingested as such with the food of the fish and stored in the liver, or what part, if any, is played by the fish in the synthesis of either vitamin-is still obscure, and reference may he made to the following memoirs ergosterol (cf. Massengale and Nussmeier, J. Biol. Chem. 1930, 87, 415, 423; Steenbock Biochem. J. 1922, 16, 518; Bills, J. Biol. Chem. and others, ibid. 1932, 97, 249; Russell, Taylor, 1927, 751; Wimpenny, Sci. Rep. Fisheries Invest. Ser. 2, vel. II, No. 5, 1929, London; [in medicine (as a eclerosing agent in, for example, Drummond, Nature, 1930, 128, 398; Gunther, the treatment of varicese veins) under the name Collin, Drummond and Hilditch J. Expt. Biol. of sodium morrhuate (cf. Haines, Lancet, 1933, 1934, 11, No. 2, 173; Copping, Biochem. J. 224, 748); the acids themselves are occasionally 1934, 28, 1516,

The statemente of Taylor and Nelson (Taylor, Iewa State Coll. J. Sci. 1931, 5, No. 4, 355; Nelson, Ind. Eng. Chem. 1927, 19, 840) that codliver oil also displays vitamin E activity is contrary to the observations of Sure (J. Biol. Chem. 1927, 74, 37, etc.) and require confirmation, especially since cod-liver oil figures as a regular constituent of satisfactory vitamm-Efree diets for experimental work (Bacharach, private comm).

According to Waterman and van Dijk (Proc. Akad. Wetensch. Amsterdam, 1931, 34, K. Akad. 1206), cod-liver oil may be hydrogenated at low temperatures under controlled conditions

without impairing the vitamin activity.

Whilst cod-liver oil itself is still largely used in medicine, its place has to some extent been taken by other fish-liver oils, cuch as hallhut-liver oils which have a much greater vitamin content and require a correspondingly smaller desage, or by mere palatable concentrates of the vitamins extracted from the Various processes for the extraction of the unsaponifiable matter from cod-hver oil for this purpose have been patented; further, high-vacuum short path distribution (" soolecular distillation") of the oil has been proposed for the separation of the active principles (cf. Waterman, van Dijk, and Imperial Chemical Industries, B.P., 452442; Rickman and East-man Kodak Co., U.S.P., 1925559).

man Acotac Co, U.S.F. 19250891, defects of the Concerning the alleged text of the Concerning the alleged text of J. Agobab. Z. Vitamunforech. 1934, 2, 09; Upenle Llakarseren Forh. 1933—4, 29, 09; 1934—5, 40, 183; Norns and Church, J. Bob. Chem. 1939, 89, 431; W.M. Coz, J.T., and Rees. Bull. Johns. Hopkins Hosp. 1934, 44, No. 6, 430; Bell, Gregory and Drumpmond, Z. Vitamunforsch. 1935, 2, 181; Thatcher, Lancet, 1936, 230, 20, L. L. Madsen, et al. Cornell Univ. Agric. Exp. Stat. Mem. 1935, No. 178; Burnek and Zimmerman,

J. Nutrition, 1937, 14, 535.

Concerping the use of cod liver oil as a healing dressing for burns and other wounds, see W. Löhr, Zentr. Chirurgie, 1934, 61, 1686; cf. Lancet, 1934, 227, 367. Steel, ibid. 1935, 229,

290; Stevenson, told. 1376.

Modern methods in the preparation of codliver oil emulsions are described by Rothenkirchen (Pharm. Ztg. 1934, 79, 804). (Concern ing the atability of vitamin A in cod liver oil emulsions, see Griffiths, Hilditch and Rae, Analyst, 1933, 58, 65).

Preparations of the scaps of the fatty acids ("British Pharmaceutical Codex"), nr nf the unsaturated acids only, of cod liver oil ace used

1 Vitamin active liver oils are also obtained from the burtot (Lots n), a freel-water fish allied to the prestigational liest, to 12, 1932; Poolsson, borsk prestigational liest, to 12, 1932; Poolsson, borsk Mag Laegyldenskap, 1931, 92, 1839, and the tunny (Zhannuz Diyanuz L.), the latter oil being especially rich in vitamin D (7 Joveknason, 7 e; behnoltz, Albera, Kgt. Jowiek Vitenska. Seisk Felt. 1939, 3,

referred to as "morrhule acid,"

(The authoris indebted to Mr. A. L. Bacharoch. M.A., F.I.C., of the Glazo Laboretories, Ltd., for helpful criticism and suggestions in cennection with this article.)

EL. CŒLESTINE. A mineral, SrSO, a seurce of atrontium.

CŒLINBLAU, CŒRULEUM v. BLUZ CERULEAN: v. COBALT.

COERULEIN, Anthracene green (v. ALI-ZARIN AND ALLIED DYESTUFFS.

CO-ENZYMES, CO-ZYMASE, Fellowing Buchner's famous discovery of an agent in the filtered liquid contents of yeast cells, capable of fermenting augar solutions in the absence of living cells, Harden end Young (Prec. Chem. Soc. 1905, 21, 189) discovered that this agent could be separated by ultra filtration into a thermosteble dislysable part-the ce zymase-and a thermolabile non-dialysable part-new called apozymase. The addition of the heiled dislysate to the mactive apozymase reatored its activity. This was in part due to the presence of inerganic bosphates and in part to an organic substance. In fact, Buchner's zymase is a mixture of many enzymes and the process of alcohelic fermentation is a complicated acries of reactions which have only recently been elucidated by Embden and by bloyerhof.

In brief, the sugar is phosphorylated before being decomposed, the primary preduct of decomposition being the phosphoric ester of a triose, probably glyccraldehyde, which by dismutation gives rise to a mixture of phospho-

glycene acid and phosphoglycenno. In the next stage the glyceric acid passes by loss of water under the influence of an enzyme into phosphepyruvic acid. At this stage the co enzyme comes into action combining with the phosphore acid group and forming free pyrivic acid The lable co-enzyme-phosphate then reacts with glucose to ferm hexosephosphate and regenerated co enzyme. The pyrivic acid as decomposed by carboxylase to acetaldehyde and carbon dioxide, the former being subsoquently reduced to ethyl alcohol. The cozymase has been identified as a complex diphosphodinucleotide containing nicetinic acid (pyridinecarboxyamule) and adenine.

In the voluntary and heart muscles, a somewhat aimilar acries of changes takes place, the essential difference being that the pyruvicacid is reduced to lactic acid. The co-enzyme is ailenylic acid, the system being

adenyi pyrophosphate

adenosine diphosphate --- adenosine-5 phosphate

(Meyerhof and Kiessling, Biochem Z. 1935, 283, 83; Lohmann and Schuster, 161d. 1935, 282, 104)

Adenylic acid was found to be a riboside 5phosphoric acid of adenine by Embden and

COFFEE. 251

Schmidt (Z. physiol. Chem. 1929, 181, 130); stitutes. For the past decade the clearances for Levene and Jacobs (Ber. 1911, 44, 746); and home consumption have been as follows: Levene and Stiller (J. Biol. Chem. 1934, 104, 299). Gulland and Holiday (J.C.S. 1936, 765) established its formula as adenine-9-riboside-5phosphoric acid,

$$\begin{array}{c|c} \mathbf{N} = \mathbf{C} \cdot \mathbf{N} \mathbf{H}_2 \\ \downarrow & \downarrow \\ \mathbf{C} \mathbf{H} \cdot \mathbf{C} - \mathbf{N} \\ \parallel & \parallel \\ \mathbf{C} - \mathbf{N} - \mathbf{C} \mathbf{H} \cdot \mathbf{C} \mathbf{H}_2 \cdot \mathbf{C} \mathbf{H} \cdot \dot{\mathbf{C}} \mathbf{H}_2 \cdot \mathbf{PO}_3 \mathbf{H}_2 \\ \downarrow & - \mathbf{O} - - - \mathbf{M} \end{array}$$

Thus the co-enzymes of alcoholic fermentation, of lactic acid dehydrogenase, of muscle and of Warburg's (Biochem. Z. 1935, 282, 1, 157) oxidation system are all closely related or identical (Adler, Euler, and Hellström, Nature,

Dec. 5th, 1936, p. 968).

Co-zymase is widely present in yeasts, in muscle, in various animal organs and tissues, particularly the liver, and in certain bacteria. It is rare in the higher plants and absent from certain races of bacteria, in which another form of sugar degradation must take place (Myrbāck and Euler, Ber. 1925, 57, 1073; Virtanen and Simola, Z. physiol. Chem. 1927,

It is obtained as a colourless, easily soluble powder (a recent summary relating to co-zymase is given hy Myrback, Ergebnisse der Enzym-

forschung, 1933, 2, 139-168).

There is evidence afforded by purification hy means of copper (Euler, Alhers and Schlenk, Z. physiol. Chem. 1935, 234, 1) or hy heating (Euler and Vestin, Z. physiol. Chem. 1935, 237, 1) that co-zymase is a molecule possessing groups whose powers of activating enzymic reactions are independent of each other. Andersson (Z. physiol. Chem. 1935, 235, 217) concludes that the substance has two independent reaction centres, one in an adenylic acid grouping concerned with phosphorylation and the other with oxidation: reduction.

E. F. A.

COFFEE. The beverage known as "coffee" is prepared by infusion of the heans or seeds of Coffea arabica, an evergreen shruh indigenous to North East Africa and Arabia. The use of coffee orginated in Ahyssinia, whence it was introduced into the neighbouring Mohammedan countries, its physiological action in promoting sleeplessness being regarded as of advantage to those attending the prolonged religious cere-monies. In spite of objections by the priests on the ground that it was an intoxicating beverage, its popularity hecame firmly established there and has continued. The use of coffee was extended through Europe during the first half of the seventeenth century and in 1652 it was introduced into England. Here it rapidly attained popular favour and during the following two centuries "coffee houses" played an important part in the social life of English cities. In 1847 the importation of coffee reached a maximum of nearly 17,000 tons, but since then it has declined, owing, no doubt, to the greatly increased use of tea and also, in some measure, to the introduction of coffee suh- follows.

Year (ended 31st Mar	rch)).			Clearances, tons.
1926-1927					15,981
1927-1928					15,897
1928-1929					15,453
1929~1930					15,308
1930~1931					16,709
1931~1932					16,340
1932~1933					15,521
1933~1934					14,782
19341935					14,763
1935~1936					15,052

These quantities may be compared with the importations into the United States, which are

over 700,000 tons per annum.

The importations into the United Kingdom are almost equally from Empire and non-Empire sources, although since 1933 the balance has been slightly in favour of the former, which in 1935-1936 reached nearly 53% of the total. This trend in favour of the Empire products (which are chiefly from British East Africa and India) is probably due to increased imperial preference under the Ottawa agreement.

The coffee tree belongs to the natural order Cinchonacea which embraces, according to Lindley and Paxton, two species, C. arabica and C. paniculata, but some botanists distinguish many other species. Four commercial types distinct from each other are now grown: (1) Arahian or Mocha coffee, having short, upright branches with a hrittle leaf, and seeds usually single in the berries. (2) Jamaica coffee, bearing longer and more pliable branches, a tougher leaf, and seeds almost always double in the herries. (3) East India or Bengal coffee, of which the leaf is smaller than that of the Jamaica variety and the berry very small. (4) Liberian coffee is taller and has larger leaves and fruit than the other varieties, but, owing to its coarse flavour, it was not until recently of much commercial use. It has, however, now attained importance, on account of its hardy hahit and power of resistance to disease, and is largely cultivated hy grafting on to its stem the more delicate and highly esteemed Arabian coffec. Trees growing wild in the Cingalese jungle attain a height of 14-18 ft., but, under cultivation and subjection to systematic pruning, the plant rarely exceeds 9 ft. and is usually less.

PRODUCTION.—Within the tropics the coffee tree thrives hest at 1,200-3,000 ft. above sea level, but grows at a height of 6,000 ft. When the mean temperature is about 70° it may grow as far north as latitude 36°. It is cultivated in the West Indies, Mexico, Central America, Brazil and Cayenne; in Western Africa, Egypt, Mozamhique and Natal; in Arabia, India, Ceylon, Siam, Sumatra, Java, Celebes and other parts of the Eastern Archipelago, Fiji and the islands of the Pacific. According to data extracted from the "International Year Book of Agricultural Statistics" the total world production of coffee for the years 1930-1935 was as

		0	Production
			. 2,363
			2.035
			. 2,330
- 1	- 1	- 1	. 2,472
	÷		. 2,078
:	: :		

Of these totals Brazil produced, on the average, about 67%. Brezil is also the greatest exporter of coffee, a preponderence clearly demonstrated hy the following data pubbshed by the Brazilian Government:

(Thousands of tons)

	1933	1934	1935	1936
Exports from Brazil— To Europe	353	344	368	337
To United States ,	486	490	489	470
To southern ports . Exports from other	68	61	75	80
countries-				
To Europe	263	294	271	323
To United States .	220	205	244	276

Although the exportation from Brazil has shown little variation, the production is restricted by the fact that the trees, after a year of beavy crops, require a season to recuperate. For commercial reasons much of the coffco produced in Brezd has been withheld from the market and to the end of 1935 over 2 million tons had been destreyed. According to data published by the Empire Marketing Board in 1932 the area under cultivation in the British Empire was about 368,000 acres compared with about 10,000,000 acres in foreign countries, whilst of the total world

exportation 3% was from Empire countries. CULTIVATION -The method of forming plantations varies somewhat in different countries not very heavy Probably the best soil is that known in Brazil as "terra roxa," a top sod of red clay 3 or 4 ft, thick with a subsoil of gravel Virgin forest land is as a rule found to be very suitable. The soil should not be very wet and steep alopes are unsuitable owing to rain wash The loss due to this cause may be repaired by good manure-well rotted dung and ground ground rock phosphate and bone meal form the reared from seeds in nurseries, whence they are transplanted to their permanent position, or they may be gathered out of the old planta-The age at which this is done varies. Thus in Brazil transplanting usually takes place when the plants are 2 years old, wholst in the West Indies the plants are about 6 months old. Fruit is produced in the fourth year and continues to be gathered for several years. The tree blossoms and bears fruit practically all the year cound and the fruit takes about four almost perpetual crop, but, as a rule, there are two principal harvest acasons in the year. Leaf beans are coasted. 20; 1934, 3). The conclusions indicate that table will indicate the variations in size :

there is no relation between the incidence of the disease and the chemical composition of the soil, and the addition of hme and fertilisers does not protect trees predisposed to attack. Whilst it is stated that Bordeaux mixture is not in itself a satisfactory remedy, certain modifications of the mixture are suggested which will prevent germination of the fungus spores. According to Sladden (Bull. Agric Congo Belge, 1932, 23, 329) infection of the harvested bean with Stephanderes hampes can be corrected by fumigation with turpentine, which does not injure the germinative power of the beans. Layere of the beans are alternated with layers of cotton cloth ansked with I ml. of turpentine per 100 sq cm. and allowed to stand in hermetically scaled containers for 3 days at 26°C.

The coffee fruit consists of a bilobed berry somewhat resembling a small elongated cherry, by which term it is described commercially, As the fruit ripens the fleshy pericarp changes colour from green through yellow and red to a deep crimson When dried, it blackens and becomes hard and brittle, resembling a shell or hush. Beneath the skin of the pericarp is a saccharino and somewhat glutinous pulp containing the seeds, which are usually double, lying face to face, but sometimes occurring singly Each seed or bean is enclosed in two membrenous coverings-an outer, consisting of tough huff-coloured tissue, technically known as " parchment," which becomes loose and readily separates on drying; and an inner film of thin white tissue, closely adherent to the bean, and known as the silver skin.

HARVESTING -The principal methods vogue for gathering and preparing the coffee berries for commerce are as follows. In Arabia. The surface soil should be good and the subsoid Egypt, and the Antilles the berry is left on the tree till completely dried, when it is easily shaken down and the dry outer envelope and the "parchment" removed by pounding in a mortar, beating with a flail, or simply rubbing in the hands. In other countries the berries are gathered by hand as soon as ripe but not dry and are spread out on the ground in layers from 4 to 6 in. deep. These are exposed to tha sun'a heat for 3 or 4 weeks with frequent bones being the best. According to Beckley sun's heat for 3 or 4 weeks with frequent (Kenya Colony Dept. Agric. Bull. 1932, 16) turnings with a spade, when the bean becomes easily removable from the fruit by gentle trimost effective fertiliser. Shelter from wind is turation. This process, however, sometimes most important. The joung plants may be imparts to the college a disagreeable flavour due to putrefactive changes in the pulp, and recourse is therefore frequently had to artificial heat. The most modern method is to "pulp" the bernes by crushing between rollers so as not to injure the beans, which are then subjected to a fermenting process in tanks, with or without the addition of water, after which they are acparated from the pulp by "washing" and dried, either by the sun or by artificial means. Coffee beans in the form of "cherry" rarely enter into commerce but frequently retain the months to ripen. There is thus in some cases an parchment and inner skin, which, however, are removed by colling and winnowing before the beans are coasted. The beans are then graded and berry fall, caused by disease, has been the late different ares by apecual machinery which aubject of considerable investigation, particularly in Kenya (Dept. Agre. Bull. 1932, 17, the ordinary flat faced beam. The following COFFEE. 253

No. of seeds in approx. 50 ml. 187 Fine hrown Java 198 Fine Mysore 203 Fine Neilgherry 203 Costa Rica . 207 Good ordinary gautemala 210 Good La Guayra Good average Santos 213 Fine long herry Mocha. 217 Good ordinary Java 223 Fine Ceylon plantation. 225 236 Good average Rio Medium plantation (Ceylon) 238 Manila 248 Ordinary Mocha 270 West African 313

ROASTING.—The object of roasting coffee is to develop its aroma and to destroy its toughness so that it may easily he reduced to powder. The process is conducted in cylinders heated hy coke fires or gas. The cylinders are usually made of strong wire gauze to permit the escape of steam and other volatile products im-mediately they are formed. Great care is necessary to ensure an equable roasting of all the heans and to attain the exact point at which the process should cease. If, on the one hand, the process is not carried far enough, the flavour of the bean is not developed and the coffee is hard to grind. On the other hand, overheating causes carbonisation and the production of a strong empyreumatic flavour, which renders the coffee infusion nauseous. The loss in weight is from 15 to 18%, of which approximately one halt is due to moisture and the rest chiefly to volatile products derived from the fat and from the sugar during caramelisation. A certain small amount of caffeine is also lost. At the same time an oily substance caffeol (q.v.) is developed, which, although present only in very small proportion, imparts to roasted coffee much of its characteristic aroma. The most favourable temperature for developing the maximum aroma during roasting is a little above 200°C. After roasting it is most important that the beans should be cooled rapidly. This is effected by spreading them on a large sieve through the meshes of which they cannot pass, hut through which air is forcibly drawn by means of a powerful exhaust connected with a closed chamber immediately below the sieve.

During the process of roasting various suhare driven off. Bernheimer (Wien Akad. 1880, 81, (2), 1032) found that, after water vapour, there is a sudden evolution of carhon distillation accompanied by dehydration and values:

polymerisation. Trigg (R.K.D. Bull. No. 3). In the interior of the hean reducing conditions obtain whilst oxidation place on the exterior. Bérnheimer (Monatsh. 1880, 1, 456) found that 50 kg. of raw coffee beans roasted in a revolving drum yielded ahout 5 litres of liquid distillate with approximately 680 g. of solid floating thereon. The filtered liquid was yellow in colour and had the fragrant odour of coffee. It was found to contain caffeine, caffeol, acetic acid, quinol, methylamine, and acetone. The solid portion was found to consist chiefly of palmitic acid. In contradiction to the above workers Hagen (Trop. Agric. 1922, 59, 269), whilst admitting that marked changes take place during normal roasting, considers that the constituents of the hean are not dissipated, there being a transformation into substances which contribute to the aroma of the roasted product. Erdmann (Ber. 1902, 35, 1846) attributed the aroma of roasted coffee to traces of acetic acid, furfural, valeric acid, an unstable nitrogenous compound, and a phenolic body resembling creosote, whilst other workers have suggested that the main odorifcrous principle is some unidentified nitrogen compound.

DETERIORATION AFTER ROASTING .- Since the essential oil rapidly escapes after the bean is broken up coffee should not be ground until required for infusion. Apart from this physical consideration, however, the bean undoubtedly gradually undergoes a process of deterioration due to chemical changes and known in the trade as "staling." Trigg (l.c.) suggests that this is due to hydrolysis and oxidation, causing alteration and volatilisation of the aromatic principles. Tunnett and Eddy (Food Ind. 1930, 2, 401) suggest that the taste of stale coffee is probably due to the oxidation of the oil. They found that staleness developed only slightly faster in ground than in unground coffee. Their work has been further developed by Bengis (Food Ind. 1935, 7, 490), who states that roasted coffee contains about 15-16% of oil and fat, much of which is unsaturated. The oil extracted from freshly roasted coffee carries in it the aroma and flavour hearing substances which are easily oxidised. On the other hand, the oil extracted from stale coffee has a rancid, disagreeable flavour. It absorbs much less oxygen which, it is therefore suggested, plays an important part in the staling.

CHEMICAL COMPOSITION .- Many investigators have published data relative to the chemical composition of coffee. Levasic (Arch. Pharm. dioxide prohably due to a process of destructive 1876, [iii], 8, 294) found the following percentage

Gummy matter.	Caffeine.	Fat.	Tannic and Caffe- tannic acids.	Celiu- lose.	Ash.	Potash.	Phos- phoric acid.
25.3	1.43	14.76	22.7	33.8	3.8	1.87	0.33
1						2.13	$0.42 \\ 0.27$
27.4	1.14	15.95	20.9	32.5	4.5		0.51
				33.0	4.9	- 1	0.46
	1				4.3	- 1	0.60
	25·3 22·6 23·8	25·3 1·43 22·6 0·64 23·8 1·53 27·4 1·14 20·6 1·18 25·8 0·88	25·3 1·43 14·76 22·6 0·64 21·79 23·8 1·53 14·87 27·4 1·14 15·95 20·6 1·18 21·12 25·8 0·88 18·80	Gummy matter. Caffeine. Fat. Caffete tannic acids. 25·3 1·43 14·76 22·7 22·6 0·64 21·79 23·1 23·8 1·53 14·87 20·9 27·4 1·14 15·95 20·9 20·6 1·18 21·12 21·1 25·8 0·88 18·80 20·7	Gummy matter. Caffelne. Fat. and Caffelne tannic acids. Cellulose. 25·3 1·43 14·76 22·7 33·8 22·6 0·64 21·79 23·1 29·9 23·8 1·53 14·87 20·9 36·0 27·4 1·14 15·95 20·9 32·5 20·6 1·18 21·12 21·1 33·0 25·8 0·88 18·80 20·7 31·9	Gummy matter. Caffelne. Fat. and Caffer tannic acids. Cellulose. Ash. 25·3 1·43 14·76 22·7 33·8 3·8 22·6 0·64 21·79 23·1 29·9 4·1 23·8 1·53 14·87 20·9 36·0 .4·0 27·4 1·14 15·95 20·9 32·5 4·5 20·6 1·18 21·12 21·1 33·0 4·9 25·8 0·88 18·80 20·7 31·9 4·3	Gummy matter. Caffeine. Fat. Caffe-tunnic acids. Cellulose. Ash. Potash. 25·3 1·43 14·76 22·7 33·8 3·8 1·87 22·6 0·64 21·79 23·1 29·9 4·1 2·13 23·8 1·53 14·87 20·9 36·0 .4·0 27·4 1·14 15·95 20·9 32·5 4·5 20·6 1·18 21·12 21·1 33·0 4·9 25·8 0·88 18·80 20·7 31·9 4·3

Commaille (Mon. Sci. 1876, [3], 6, 779) found | should be taken and first extracted with other. in undressed Mysore roffee :

						**
Fat						12 68
Glucose						2 60
Legumin	2.00	in		٠	٠	1.52
Albumen				٠	٠	1 04
Caffeine	٠	٠	٠	-	٠	0 42-1 31
Ash						3 88

Five different commercial types of Ceylon coffee gave the following results published by the Imperial Institute :

		Percentage	
	Mia	Max	Average
Moisture	10 1	106	103
Caffeine	21	24	23
Crude proteins .	11.2	13 4	125
Fat	61	78	69
Carbohydrates .	493	52 1	50 7
Crude fibre	13 5	153	14 4
Ash	33	35	34

It will be observed that these eamples were onusually rich in casseins whilst the proportion of fat was low. Data published by the same authority for rosseo grown in Sierra Leone were more normal, approximating closely to the values for Arabian coffee, 112, mosture 82, caffeine 1-3, crude proteins 85, fat 10 1, nitrogen-free extract 50 6, crude fibro 17 8, and as 35%. Valenzuele (Philippine J Sci. 1029, 35%. Valenzuele (Philippine 40, 349) gave the average percentage composition of acven varieties grown in the Philippines as follows: moisture 986, caffeine 187, fet 8 71, reducing augar 5 22, crude fibro 19 65, nitrogenous substances 12 95, and ash 4 10.

Caffeine. Of the substances which have been isolated from coffee that which has attracted most attention is caffeine. It is interesting that the two other popular beverages, tea and cocoa, rontain respectively the alkaloids theine, identical with caffeine, and theobromine, which la closely related to it. It has been soggested that the caffeme is present ac caffetannete, whilst Gorter (Annalen, 1908, 358, 327) states that the caffeine occurs in coffee combined with potassium as potassium-caffeine chlorogeuste,

C, H,,O,K(C,H,,N,O,),.

Calleina also vecus in many other commodities, esfler, is unextrain in its competition and very and the method of its determination has there- susceptible to change. According to Gorter fore received considerable attention. The per- (Annalen, 1908, 358, 327) it is associated with centago of cassence in collec, and particularly in the so called de-tassensed cosses, is so low, however, that some methods suitable for general 4 3% in the rosated cosses (Hoepita, Z. Uniters. application are of little value in this connection | Lebessim 1033, 68, 233). The same worker and only those especially devised for coffee need (Chem. Zig. 1932, 56, 901) states that in the be mentioned. Lendrich and Nottholm (Z pursers, Nahr.-Geaussim 1909, 17, 241) propose to be extraction with carbon ictrachlorule, and NaOH. Ilis method has been modified by transferrace to solid parsfirm which is then Pilicker and Keilholz (Z. Unters, Lebesum 1934, extracted with water. After cleaving with 68, 971. Jurany (Z. anal. Chem. 1933, 94, 225) proposes a method involving extraction with exported to dryners, the residue extracted with alcohol and transfer through solid parafiln to an electronic middle, and weached. Fradler and annous soliton which is clearly with the state of the control of the c application are of little value in this connection Lebensm 1033, 68, 238). The same worker chloroform, dried, and weighed. Fendler and aqueous solution which is cleared with lead Stuber (Z. Unters. Nabr. Genussm. 1914, 23, 9) acctate, the optical rotation then giving a whist confirming this method suggest explain measure of the chlorogenic acid present. wants confirming this method suggest efficient measure of the chlorogenic acid present, modifications which render the estimation baser. According to Jurany samples of raw collect were constituted in the contract of the co expeditious. Vautier (Mitt. Lebensm. Hyg. 1919. found by this method to contain 8-9% of chloro-19, 272) suggests that at least 10 g. of the sample | genic acid, q v.

The crude coffeine should then be extracted with chloroform, evaporated to dryness with sodium carbonate, again extracted with chloroform five or aix times, the extracted caffeine thea being dried and weighed. Bonifazi (Mitt. Lebensm. Hyg. 1926, 17, 307) confirms Vautier's method. Various modifications have been suggested. Thus Ugarte (Chemia, 1930, 7, 490) proposed that before extraction the material should first be carbonised cautiously in a Kjeldahl flask, care being taken that none of the white vapour formed is allowed to escape. Bonn ood Desgrez (Aon. Falsif 1931, 24, 546) propose that after extraction the residue from evaporation should be taken up in 10 ml of 5% HCl and an excess of edicatungstic acid solution added. heating and prolonged standing the precipitate is filtered off, ignited and weighed, the weight of the precipitate multiplied by 0 2236 representing the hydrated caffeine. Helberg (Mitt. Lebensm. Hyg 1933, 24, 54) states that best results are ohtsmed if the powdered coffee is first rubbed with a mixture of lime and quartz cand. Herndlhofer (Mikrochem. 1932, 12, 227) proposed an interesting rephelometric method for measuring the turbidity produced by adding sodium phosphotungsteic to a cassence solution. The amount of cassens in "decassensed" rostro varies considerably according to the netoro of the original coffee and the process employed. Guillaume and Lefranc (Bull Su Pharmscol. 1935, 42, 14) state that the quantity of soluble substanco in " caffeine-free " coffee is only twothirds of thet in the untreated material and the caffeine may be as low as 0 0004%. Braunadorf (Z Unters. Lebenam 1933, 65, 460) found that in an infusion of 8-11 g. in 150 ml. " caffeine poor " coffee gare up to 22 mg. of caffeine, " caffeine-free" coffee gare up to 8 8 mg., and coffee gare up to 72 mg. Bonifazi (Mitt. Lebenam. 11yg. 1926, 17, 307) found that 20 g. of decaffeinised coffee usually contains about 8 4 mg. of caffeine. For the estimation Borgmann (Apoth. Ztg. 1931, 71, 56) proposes a method involving Soxhlet extraction, whilst Chifford (J. Assoc. Off, Agric. Chem. 1931, 14, 553) supports a method based unon sublimation

Coffetannie acid, which is said to be present in

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Chem. 1895, 8, 469) state that sugar occurs in the raw bean only as a glucoside. This is in contradiction to the findings of other investigators. Thus König and Bell found from 5 to 9% in raw coffee. In support of this finding Bell points out that when the saccharine extract of the heans is boiled with a little dilute sulphuric acid a quantity of glucose is obtained nearly equivalent to the weight of dry extract operated upon, a result inconsistent with the presence of sugar as a glucoside. Bell considers that this sugar is not present as sucrose, but the chemists of the United States Department of Agriculture (Bulletin 13) have isolated cane sugar in considerable amount as pure, well-defined crystals. It is usually accompanied by some invert sugar probably formed during the process of extraction. It is generally admitted that there is practically no sugar in roasted coffee, that present in the raw bean being nearly all caramelised during roasting.

Cellulose.—In coffee this is very hard and very difficult to define under the microscope, especially after roasting, although the chief characteristics of the cellular tissue are more or less retained. It is dissolved hy an ammoniacal solution of copper oxide and gives a hlue colour with sulphuric acid and iodine. Remnants of the "silver skin" are always present even after roasting, and its structure, a thin memhrane with adherent thick-walled, spindleshaped and hollow cells, is quite peculiar to coffee

and a useful means of identification.

Fat v. COFFEE BEAN OIL.

Protein, chiefly in the form of insoluble alhuminoids, occurs in coffee, both raw and roasted, to the extent of from 10 to 14%, the soluble alhuminoids constituting about 2%.

Mineral Matter.—The total amount of ash varies from 3.5 to 4.0% in raw, and from 4.0 to 5.0% in roasted coffee. The following percentage compositions are given by the United States Department of Agriculture (Div. of Chem., Bull. 13, 904).

		Mara-		
	Mocha.	calbo.	Java.	Rio.
Sand	1.44	0.72	0.74	1.34
Silica (SiO ₂)	0.88	0.88	0.91	0.69
Ferric oxide (Fe ₂ O ₃)	0.89	0.89	1.16	1.77
Lime (CaO)	7.18	5.06	4.84	4.94
Magnesia (MgO)	10.68	11.30	11.35	10.60
Potash (K ₂ O)	59.84	61.82	62.08	63.60
Soda (Na ₂ O)	0.48	0.44		0.17
Phosphoric acid				
$(P_2 O_5)$	12.93	13.20	14.09	11.53
Sulphuric acid				
(ŜO ₃)	4.43	5.10	4.10	4.88
Chlorine	1.25	0.59	0.73	0.48
		_		-

Ludwig (Arch. Pharm. 1872, [iii], 1, 482) states that the silica increases from the root upwards to the leaves and pericarp of the fruit, after which it declines in the parchment skin and is nearly absent in the bean.

Physiological.—The primary value of coffee as an article of human consumption is its stimulative property. Indeed, it can, when taken in a strong infusion, have such an exhilarating effect that, as already stated, attempts were (Z. Nahr. Genussm. 1900, 3, 681) describes the

Sugar.—Herfeldt and Stutzer (Z. angew. made to prevent its use in Mohammedan countries hem. 1895, 8, 469) state that sugar occurs in the on the ground that it was an intoxicant.

According to Lehmann and Weil (Arch. hyg. 1923, 92, 58) the action of coffee on the nervous system is to be attributed to the caffeine present. For the same reason coffee is a stimulant to gastric secretion (Haneborg, Acta. Med. Scand. 1924, 61, 228; and Farrell, North Western Univ. Amer. J. Physiol. 1928, 85, 672) although Bandoin (Bull. Soc. hyg. aliment, 1920, 11, 655) found that coffee has a decided retarding effect on peptic and pancreatic digestions which is proportionate to the tannin content. A strong infusion of coffee has a distinct diuretic effect which, probably owing to the action of lecithin, can be inhibited by the addition of milk or (more effectively) cream, Hitzenberger and Roller (Wien. Arch. inn. Med. 1935, 27, 133). Pietro di Mattei (Arcb. Farm. sperim. 1923, 35, 5) found by experiments on pigeons that roasted coffee had a marked curative effect in cases of polyneuritis, although raw coffee and pure caffeine were without effect.

ADULTERATION.—Although the adulteration of coffee in this country is now practically nonexistent, it is very common abroad and the number and variety of the adulterants are almost unlimited. Many of these are hest detected by means of the microscope. Digestion with water affords a ready means of differentiating hetween the pale soft-walled tissues of the cereals and roots, often used as adulterants, and the darkcoloured and hard-walled tissue of coffee. The cereals may readily he identified hy their characteristic starch granules. The chemical differences hetween coffee and its adulterants aro also strongly marked, as the sophisticated products usually show high aqueous extract and sugar and low fat and caffeine contents. The density is also a useful factor in the preliminary examination, that of raw coffee varying hetween 1.041 and 1.368 and that of roasted coffee between 0.500 and 0.635. In the Paris Municipal Lahoratory, Regnault's volumenometer, as Lahoratory, Regnault's volumenometer, as modified by Dupré, is used for this purpose. Colouring matters are frequently employed to disguise damaged beans and give them the appearance of superior quality. Among the substances so used are chrome yellow, yellow ochre, indigo, Prussian-blue, azo-dyes, and soapstone for the raw bean; and burnt umher, oxide of iron, and Venetian red for the roasted bean. The inorganic colouring matters are best sought in the ash or in the sediment obtained by sbaking the beans in cold water; indigo and Prussian blue by their usual reactions and other colouring matters by extraction with alcohol. Glazing is a process resorted to for the alleged purpose of preserving the roasted beans. The substances used, a mixture of egg albumen with glucose or gum, are said to have the effect of clarifying the infusion. The improved appearance of the beans and increased weight due to the retention of moisture, are probably also important considerations. The latter result is sometimes attained by the addition of sugar before roasting or by steaming the beans and coating them with glycerine, palm oil or vaseline to prevent evaporation of moisture. Bertarelli

256 COFFEE.

adulteration of roasted coffee beans with a solution of borax, which not only gives the beans a glossy appearance but also adds as much as 12% to their weight. Shellac is occasionally used for glazing, and Nottbohm and Koch (Z. Nahr. Genussm. 1911, 21, 288) draw attention to the fact that shellac is sometimes coloured hy arsenic trisulphide. Coffee beens which had been glazed with arsenical shellar were found to be atrongly contaminated with arseme. Zur-briggen and Scheurer (Mitt. Lebensm. Hyg. 1931, 22, 377) found that many substances used as coatings for coffee can be identified by washing the grains with a suitable solvent, evaporating and comparing the fluorescence of the extract under a mercury lamp with that of pure specimens of the aubstances suspected

Although the addition of chicory (q.v.) to coffee is often regarded as a form of adulteration, public preference for this admixture justifies the view that chicory is an adjunct to, rather than an adulterant of, coffee if its presence is disclosed at the time of sale. It would appear, however, that chicory is declining in favour, the consumption in the United Kingdom having steadily fallen from 3,200 tona in 1927 to 700 tons in 1936. Chicory has little in common with coffee except a considerable proportion of augar which is converted into caramel in the process of roasting and thus imparts to the infusion a deep colour, a hitter taste, and an acoma somewhat resembling that of coffee. Smith and Bartell (Univ. of Mich. Dept. of Eng. Recearch, Sept. 1924) show that, owing apparently to a process of chemical reaction between the caffeol of the coffee and the soluble constituents of the chicory, the acomatic caffeel is held in the infusion. An elaborate research i into the natuce of coffee and chicory with a view to the detection of adulteration was made at the instance of the Commissioners of Inland Revenue in 1852 by some of the leading botanists and chemists of the dey, and the results aco of great practical value even at the present time. The following table, prepared during this investigation, shows the ap gr. at 60°F, of colutions made by treating I part of each of the aubatances named (roasted and finely ground) with 10 parts of water (hy weight), raising the mixture and cooling.

English chicory .		. 1021-7	
Dandelion root .		. 1021-9	
Red beet		. 1022 1	
Foreign chicery		. 1022 6	
Gnernsey chicory		. 1023 2	
Mangold wurzel		. 1023 5	
Roasted figs .		. 1024 9	
Matze		. 1025 3	
Bread raspings .		. 1026 3	

It will be seen from the above table that the low specific gravity of the coffee infusion distinguishes it from roots and ecreals, and when chicory is the only aubstance admixed with the coffee its percentage may be calculated with approximate occuracy by a comparison of the gravity of the infusion of the mixture with the average gravities of coffee end chicery given in the table. As an alternative method Savini (Annah Chim. Appl 1923, 13, 217) auggests boiling 10 g. of the sample for 5 minutes with 200 ml of water. The infusion is then cooled. made up to 250 ml, and filtered; 25 ml, of the filtrate are evaporated to dryness and the residue weighed. Variation from the normal figuee for coffee indicates some form of adulteration. Bunco and Mostra (Analyst, 1932, 57, 708) state that the average extrect obtained from Indian coffee is 28%, which is about 4% higher then is ordinarily assumed for coffee, and that therefore the origin of the coffce must be taken into account in applying the above methods The relative colouring powers of reasted coffco and chicory afford a further simple test for the presence of chicory or other substances containing caramel. A few grains of the sample are placed on the aurface of cold water in a glass vessel, without stirring, when the particles of foreign aubstance each emit o heownish-coloured eloud, which is rapidly diffused in streaks through the weter. Pure coffee affords no appreciable coloration under annilar conditions. The chicory readily sinks, whilst nearly the whole of the coffee remains at the auriace. If examined under the microscope the sediment of chicory which reaches the bottom of the vessel will be found to exhibit the characteristic large cells with dotted vessels and branching lactifecous ducts. Even to the touch of a needle there to boiling point for half a minute, then filtering is a difference, for whilst chicory under these conditions will feel aoft, the coffee grains remain hard. Winter-Blyth ("Fooda; Their Com-position and Analysis," p. 368) thus summarises the influence of chicery on the coffee mixture : the respective proportions of gnm, fatty matter, tannio and caffetannic acids, and caffeine are decreased, the proportion of augar is increased, and the constitution of the ash is profoundly modified, especially in the proportion of ailica, which is not found to any great extent in coffee sah. In amphication of the last point the following may be eccepted as representing the principal differences in the composition of the respective ashes :

	Coffee.	Chicory.
Silica and aand .	_	10-09-35 85
Carbonic acid	14 92	1-78-3 19
Senguioxide of iron	0 44-0 98	3 13-5 32
Chlorine	0 26-1-11	3 28-4 93

i

WAX.

There is a marked difference in the proportion of ash soluble in water, that from coffee heing about 75-85% and from chicory 21-35%.

"Decaffeineised" Coffee.—The public demand for "decaffeineised" coffee has resulted in considerable search for a method which, while removing some or all of the caffeine, will yield a product possessing unimpaired the flavour and aroma of the original bean. Many of the processes thus devised have heen patented and it is possible that, at least in some cases, commercial considerations have, in part, formed the hasis for the favourable claims advanced on their hehalf. In any case the so-called "decaffeineised" coffee almost invariably contains some caffeine which may vary from a trace up to 1%. The basic principle underlying nearly all the processes is some form of extraction hut the solvents used and the treatment of the beans show wide variation. When an aqueous extraction is relied upon the water may he untreated, rendered alkaline, or loaded hy the solution of sugar or other substances, in order to effect the selective removal of the caffeine from the beans. The organic solvents used include benzene, toluene, ketones, alcohols and chloroform. In some cases solvents are used in rotation whilst in others an emulsion of chloroform or henzene with water is used. The extraction sometimes takes place under pressure whilst in other methods the beans are crushed, rolled, or steam-heated before or during the extraction. It is a common practice to remove the last traces of the solvent by passing a current of hot air or steam over the beans at the end of the process. In some cases no attempt is made to conduct a selective extraction. Conditions are imposed which result in substances other than caffeine being removed. The solution is subsequently treated with a solvent in which the caffeine is readily soluble and the residual extract returned to the beans by spraying before roasting. As an alternative to extraction by treatment of the beans with a solvent it is claimed that infusions of coffee can be "decaffeineised" by passing through charcoal or a layer of silica gel.

SUBSTITUTES OR IMITATIONS, although not very common in the United Kingdom, are more prevalent in the coffee-drinking countries of Europe and America. They are prepared from a variety of different substances. In addition to chicory, these include acorns, heet refuse, hran, cereals (barley, maize, malt, cats, rye and wheat) caramel, carrots, dandelion root, figs, lupin seeds, mangolds, parsnips, peas, heans, coffee husks, cassia seeds, date stones, etc. In general these may be identified by the methods proposed ahove. These preparations are sometimes offered quite openly as substitutes, hut imitation heans have heen prepared hy mixing any of the ahove with gum, molasses or other adhesive material and skilfully moulding the mixture into the form of coffee berries. The following are some of the imitations or sub-

stitutes which have heen placed on the market:

Negro or Mogdad coffee made from the seeds
of Cassia occidentalis.

Massaendar coffee from the seeds of Gærtnera vaginata.

Vol. III.—17

Kunst Kaffee from roasted roots or cereals.

Pelotas coffee and Coffee surrogate from ground

Date coffee from date stones or dried dates mixed with a small proportion of coffee.

"French Coffee" is a description sometimes applied to a mixture of coffee with burnt sugar and a large proportion of chicory. As these ingredients readily absorb moisture on exposure to the air this mixture is usually preserved in tins.

F. G. H. T.

OIL

and

BEAN

COFFEE

Apart from a waxy coating on the outer surface of the hean, coffee heans contain from 5-12% of a fatty oil; the wax-free oil has an iodine value of about 100, a hydroxyl value ahout 23, and saponification value ahout 195, and contains ahout 12.6% of unsaponifiable matter (Schuette, Cowley and Chang, J. Amer. Chem. Soc. 1934, 56, 2085; cf. Meyer and Eckert, Monatsh. 1910, 31, 1227). The fatty acids consist chiefly of palmitic, linolic, and oleic acids, with smaller amounts of myristic, stearic, and arachidic acids; higher fatty acids (tetracosoic acid, the "carnauhic" acid of some writers) are present in the wax from the outside of the bean and are, therefore, found when the total ether-extract of the entire bean (saponification value 150-195, iodine value 80-100) is examined (cf. von Noël, Pharm. Zentr. 1929, 70, 69; Heiduschka and Kuhn, J. pr. Chem. 1934, [ii], 139, 269; Bengis and Anderson, J. Biol. Chem. 1934, 105, 139; Bauer and Neu, Fette u. Seifen, 1938, 45, 229). A phytosterol, m.p. 138-139°, and a substance, C₁₉H₂₆O₃ (m.p. 143-143·5°C.), termed "kahweol" (from the Arahic "qahwch," C₁₉H₂₆O_{3,}, kahweol', meaning coffee) have been separated from the unsaponifiable matter (Bengis and Anderson, J. Biol. Chem. 1932, 97, 99; Dillingham and

Thompson, Univ. Hawaii, Occas. Papers, 1934, No. 19; Bauer and Neu, l.c.). The oil is also stated to possess antirachitic activity (cf. von Noël and Dannmeyer, Strahlentherap. 1929, 32, 769), which, like the characteristics of the oil, is not affected by the roasting of the coffee prior to extraction of the oil. Coffee oil docs not appear to possess much technical interest at present, although a certain amount of the crude fat (including the wax) can be recovered as a hy-product from the extracts ohtained in the decaffeising of coffee hy the solvent process; the recovery of the oil from coffee scheduled for destruction in Brazil has also heen proposed. It is stated that during the War of 1914—1918 used coffee grounds were collected in Austria for the sake of the 12% of oil which was extracted therefrom (for soap-making?) (cf. Fahrion, Z. angew. Chem. 1917, 30, 125; Munk,

COGNAC v. BRANDY.

"COGWHEEL ORE" v. BOURNONITE.

E. L.

Allgem. Oel. u. Fett-Ztg. 1932, 29, 13), a

practice which has recently heen revived in Germany.

COHUNE NUT. The fruit of the cohune palm (Attalea Cohune Mart.), grown largely in Honduras. The nut is very similar in structure to the coconut, hut is smaller. Its primary use

is for the production of oil utilised largely or cordage. The short fibre or tow is used in the for soap-boiling. The fleeh of the nuts contains form of curled fibre as a substitute for horse hair to sop-coung. I meen out and not contains from of certed three as a substitute for force hair 4%, of water, 63-71%, of oil, and approximately as a stuffing material in poblestery, and the dust 20% of protein. The principal protein is a and refuse as a manure and for other gardening globulin which may be extracted with hims purposes.

And precepitated by addition of ammonium Cours composed of strong, elastic, reddishand precepitated by addition of ammonium. sulphate. Jones and Gersdorff give the nitrogen distribution of the globulin as:

					70
Amide N .					7-5
Humin-N .					09
Cystme-N .					0.5
Arginine-N					308
Histidine-N					26
Lysine-N .					79
Amine-N of	filtr	ato		i	47.9
Non-amino-N			nte.	i	2.3

Tryptophan (0.65%) has also been detected (J. Biol. Chem. 1920, 45, 57; 1924, 62, 183) A. G. Po. COHUNE NUT OIL v. COCONDT OIL

GROUP. (this vol. p. 243d.) COIR (or coconut fibre) is derived from the fibrous husk (mesocerp) of the fruit of the coconut pelm (Cocos nucifera Linn). This palm is distributed throughout the coastal regions of the tropics. It is cultivated extensively in India, Ceylon and Malaya, and is largely grown in the Philippine Islands and in South American countries. Some varieties of the pelm are said to yield fibre of a finer quality then others; the quality is also influenced by the situation of the plantation, the fibre yielded hy palms growing near the see being finer than

The fibrous meterial in its rew state consists of a mass of fibre strands of varying length associated with corky and other non-fibrous as the nuts ripen the fibre becomes coarser and a longer period of steeping or retting is necessary, with the result that the cour ecquires a dark

that from inlend trees.

colour. In the preparation of coir by the native methods the husks are detached by striking contaming bracksh water, or in cages or wickerof the husks is effected by machinery, and the steeping is carried out in steam heated tanks. machine, the hask is passed into the extractor or breaking-down machine, in which it is comqualities. The coarser and stiffer or "brush"

brown filaments which are harsh to the touch and rather brittle These filaments are thick m the middle and gradually taper towards the ends, the diameter in the widest part varying from 005 to 03 mm.; they ere frem 15 to 32 cm. long, and either round or elliptical in cross-section. The fibre is very resistant to the action of water and is so light as to float on it. and it is therefore of special service for the mannfacture of ships' ropes.

The filaments are composed of fibre-vascular tissue consisting chiefly of very chort, irregularly thickened fibree of uneven diameter; these nitimate fibres are about 0 4-1-0 mm, long and 12-20 μ in dismeter.

The fibre substance consists of a highly lignified form of cellulose On microchemical treatment with iodino and sulphuric acid, it gives a golden yellow colour, and with aniline sulphate an intense yellow colour; it is not ettecked hy Schweitzer's reagent

COKE MANUFACTURE AND THE RECOVERY OF BY-PRODUCTS.

INTRODUCTION .- Previous to the nmeties, practically the whole of British metal lurgical coke was made in the old heehive oven, which, as will be seen from what follows, is e semi-combination process, involving the loss of the whole of the gas and by products, together with a portion of the possible coke yield. With the gradual disappearance of the old prejudices against coke made in by-product ovens, the tissue. The best fibre is obtained from fruits latter end much more economical process begen that are gathered before they are quite ripe; to assert itself, and in 1898, 1.25 million tons of coal were being cerbonised in hy product coke ovens; in 1905 the quantity of coel so treated reached 3 31 million and in 1935 about I7 million tons. Meenwhile, the number of beehive ovens in operation had decreased from methods the husks are detached by striking 16,037 in 1910 to 592 in 1935. According to the fruits on sharp spikes fixed firmly in the figures published by the Coal Conservation ground. The husks are immersed in pits Committee at the end of 1917, some 8,700 by product overs were in operation in Great work enclosures in backwaters, and left for Britain (of which 8,000 were fitted with benzol several months. On modern estates the removal recovery arrangements) with a total carbonising capacity of 17-5 million tons of coal per annum. In 1917, 166,354 tons of ammonium sulphate When sufficiently softened, the husks are were produced in British coke oven plants. The withdrawn from the water and are then beaten practice of by product recovery had become by hand with wooden mallets or passed through a universal in Germany years ago; whilst in the crushing machine in order to free the fibres from United States the by-product planta, either the corky tissue. After leaving the crushing building or in operation et the end of 1917, were espable of producing 24 million tone of pletaly distinct measures in wares is come come per amum. In view of this expladeption pletaly distinct product is then of the hy-product coking process in all these dried and treated by a "willowing," machine countries, the following article will be confined to remove dust end other chefrs. The fiver to a description of modern coking methods is usually sorted by a combing or backing viewed chefry from a by-product analpoint, process into grades of different lengths and with only a passing reference to the wasteful coke per a mum. In view of this espid adoption bechive process.

fibre is used as bristles for brush-making, Illistory of the By-Product Cokino Pro-whilst the longer and finer " mat " or spinning cress — The earliest reference which we can find HISTORY OF THE BY-PRODUCT CORING PROfibre is employed for the manufacture of matting of any observation being made on coal as a source of hy-products is that contained in the memoirs of Johann Joachim Becher, dating about the end of the seventeenth century; Becher states that he has found means for treating coal "so that it no longer smoakes nor stinks," and in doing so made a tar equal to the Swedisb.

The use of coke in iron smelting hecame general within 50 years of Becher's discovery, and a French metallurgist, De Gensanne, descrihes a process working at Sulzhach near Saarbrucken prior to 1768, where coal was coked for iron smelting, tar heing recovered as a hy-product; Stahl is said to have been the inventor of this process, and be produced there-from an "oil," hitumen, sal-ammoniac, and lamp-hlack.

In 1781 Archihald, Earl of Dundonald, was granted a patent for "allowing coals to hurn or ignite without flaming, so by their own heat to throw off the tar or oils that they contain. Dundonald stated that the only method wherehy tars could he obtained prior to his discovery was hy a distillation in closed retorts, where the admission of external air was prevented and where other coals were required for heating hesides the coal contained in the closed vessel. It is interesting to note that Dundonald was well aware of distillation in closed retorts, which is the basis of modern hy-product coking methods, and it is also evident that his knowledge of closed retorts was 16 years previous to the introduction of coal gas for street and household illumination in 1797.

In 1838 Frederick Neville obtained protection for an invention which consisted in beating ordinary gas retorts hy the waste heat from beehive coke ovens, the distilled gases from the retorts being passed through condensing and scrubbing appliances. William Newton, in 1852, introduced the use of an aspirator or pump for withdrawing the gases from coke ovens through condensers and scrubbers. Newton likewise mentions the use of acids for removing the ammonia from the gases. In 1860 W. H. Kingston patented a coke oven provided with external furnace, from which the heat evolved hy the comhustion of an auxiliary supply of coal was conducted round the oven through a system of flues; the cvolved gases were passed through a series of pipes and condensing vessels for the recovery of the by-products. Kingston's patent is especially noteworthy, since he recognises that the true method for the recovery of hy-products from coal lies in heating the coal in closed chambers, and not in chambers to which air is admitted.

With regard to Continental practice, in 1850 Pauwels and Duhouchet, and in 1862, Pernelot, introduced systems of carbonising coking coals in long rectangular chambers, heated from helow by the regulated comhustion of part of the gas in flues specially constructed under the floor, for the purposes of manufacturing metallurgical coke and removing tars, oils and ammonia. The Pernelot system was tried in the north of England in 1870, hut the coke produced was too soft for metallurgical purposes.

plant at Commentry, on the closed-retort system with bottom flues, but made no attempt to recover by-products. Later Knah was joined who (1866–1873) introduced Carves, improved methods of heating hy side flues in addition to the hottom flues, thus obtaining a more even distribution of temperature and a quicker rate of carhonisation. Carvès likewise utilised the gases (from which the by-products had heen extracted) in the side-wall flues, in addition to the auxiliary coal as employed hy Knah. The Knab-Carves ovens at Commentry gave the following results:

							70
Large	cok	е					70.00
Breeze							1.50
Dust							2.50
Grapb	ite						0.50
Tar							4.00
Ammo	nia	cal	liat	ıor			9.00
Gas							10.58
Loss							1.92
							100.00
							100.00

Further plant on the Carvès system were erected at Bessèges and Terre-Noire, near St. Etienne, between the years 1866 and 1873, and were still further improved; the Carvès ovens working at Besseges in 1880 were huilt with carbonising chambers only 24 in. wide, to facilitate rapid beat penetration and quick carbonisation. In 1881 H. Simon of Manchester introduced a recuperator, whereby the air necessary for the combustion of the gas in the side-wall flues of the Carvès oven was prebeated by the waste gases escaping from the flues, and thus the rate of carbonisation was still further increased, the combined work of the two investigators resulting in the wellknown Simon-Carvès oven, which may be beld to be the precursor of the modern by-product coking retort.

Following on the success of the Simon-Carvès process, other coke-oven patents were filed, viz. Semet-Solvay, 1880; Hüssener, 1881; Lürmann, 1882; Otto and Brunck, 1883; but although development was rapid on the Continent, little was done in this country before the nineties, owing to the prejudice of iron smelters against hy-product coke; that this prejudice has now disappeared will be seen from the rapid adoption of the process, indicated hy the figures at the beginning of this article. The hy-product coking plant which have generally found acceptance in this country are the Otto-Hilgenstock, Simon-Carvès, Koppers, Semet-Solvay, Simplex, Coppée, and Hüssener.

Non-By-Product and By-Product Ovens COMPARED.—Prior to the establishment of the hy-product coking process, the type of oven universally employed was that known as the heehive. This oven (Fig. 1) consists of a fireclay chamher, heehive shape in elevation and circular in plan, provided with a loose hrick charging door at ground level, and with an exit flue in its crown for carrying the waste gases to the chimney. The coal is charged into the oven by spades, through the charging door, to a height of In 1856 Knah erected a coal-distillation 2 ft. 6 in. to 3 ft., is then ignited, and the 260

charging door bricked up and plastered, a small | hole 6 in. square being left for the admission of air. The upper layer of the coal hurns and sets up distillation of the under layers, the evolved gases passing upwards and hurning in the dome of the oven, with the air drawn in The dome is thus acon heated at the door. up to a high temperature, end by radiation powerfully assists in carbonising the charge of coal below. This operation of distillation and immediate combustion proceeds until the whole of the volatile constituents of the coal have been evolved, which for a 10 ton charge occupies 70 hours. The coke is then quenched with

water eed withdrawn by rakes. It will be seen that by this process, the whole of the volatile products, that is to say, volatile hydrocarbons and ammonia together with the permanent gases, are burned ammediately they are evolved within the coking chambers itself : further that it is not possible so to regulate the combustion of these gasee without burning a portion of the coke in addition, in ectual practice, from 15 to 20% of the coke is hurned to waste. For the reason, too, that the coke is quenched inside the oven, the temperature of the latter is lowered, so that before the next charge can be coked, considerable expenditure of heat is necessary to raise the oven to coking temperature. The removal of the coke hy hand rakes is also tedious and expensive

With the object of overcoming the main drawbacks of the beebive oven, Coppee introduced a non by product retort of rectangular form, from which the coke could be discharged en bloc hy a ramming machine and quenched outside the oven, thus expediting the operation oven (Fig. 2) the gases evolved from the coal are drawn into flues in the side walls where they ere burned, and in this way com-bustion within the coking chember and loss of coke, as occur in the beebive oven, is e voided.

The by product retort (Fig. 3) is hult on the seme lines as the Coppée, that is to say, it is e rectangular chamber heated by fines in the side walls, and the coke is discharged by a ramming machine; the essential point of difference hes in the fact that in the Coppée the whole of the gases containing the hy-products are drawn immediately they are evolved into the side wall heating flues, where they are hurned; in the hy-product retort, the gases and products are drawn from the oven through cooling and washing plant for the recovery of the tars, ammonia, and benzol, and the permanent gas only is burned in the flues after the said products bave been extracted. In the Coppee, the whole of the permeoent gas is hurned in the heating flues ; in the hy-product oven, only one-half to two thirds of the gas is utilised in the flues, the remainder being available for beating or power purposes.

The diagrams (Figs. 1, 2, and 3) clearly

types of oven discussed above.

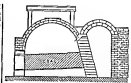
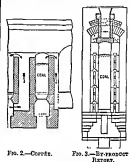


FIG. 1.-BERRIVE.



by products and a considerable yield of power or heating gas. The average yields (calculated upon 100 parts of dry coal) of coke and by-products obtainable from British coking coals of different origin by the most up to date methods, ere approximately es follows:

Origin Durham South Yorkshire	Coke	Anhyd tar.		Ammonium suiphate.
Durham	75 0	3 85	0.93	1.16
South Yorkshire	700	3 62	1.37	1 35
Lancashire	70 O	3 50	1.20	1.25
South Wales . Scotland	800	2 62	0.60	0 95
Scotland	68 0	3 25	0.70	1.00

COME OVEN DESIGN AND CONSTRUCTION. A modern by product coke oven is a rectangular chamber from 40 to 45 ft. loeg, 12 to 20 ft. high, and from 14 to 20 ins. wide. It is closed at each end by a cast iron or steel door with firebrack lining, which is lifted by a small crane whenever the contents of the oven are to be discharged. There are three or four holes in the roof through which the coal is illustrate the essential differences in the three charged into the oven from an electricallydriven charging lorry which runs on rails along "It will be seen, therefore, that in addition to the battery of evens. At one or both ends of obtaining the theoretical coke yield, and a large the oven there is an iron" ascension pipe," output, the by-product retort yields valuable through which the gas and volatile matter period are led into the "hydraulic main" running along the whole battery of ovens, much

as in gas-works practice.

The ovens and heating flues are usually constructed of silica brick, consisting of silica 92 to 96%, alumina 2 to 3%, lime 1.3 to 2.5%; such bricks are more stable at high temperatures and have a higher heat conductivity than the more aluminous bricks formerly employed; consequently, higher flue temperatures can be carried, as a result of which the speed of coking is increased while the heat economy of the system is improved, leading to a decrease in the quantity of gas required to heat the ovens.

The charge of such an oven is from 15 to 25 tons, and the coking time from 12 to 22 hours, equivalent to a throughput of from 25 to 30

tons of coal per day.

Coke ovens are built in batteries of from 25 to 160 units; they are heated by gas which is burnt in flues between the ovens so that the charge is carbonised by being heated from either side. The design of the heating flues varies much, but practically all modern coke ovens have vertical heating flues, the underlying principles being to economise as far as possible the amount of gas required to effect the carbonisation, and to enable the greatest possible arca of oven wall to be uniformly heated. The gas used for heating may be either part of the coke-oven gas after it has been passed through the by-product plant for recovery of tar, ammonia, benzol, etc., or provision may be made for heating the ovens by low-grade fuels such as producer or blast-furnace gas. In any case, however, the air required for combustion is pre-heated before it enters the combustion flues at the expense of part of the sensible heat in the burnt gases passing from the ovens. Where producer or blast-furnace gas is used for heating, such gas is also pre-heated in the same way as the air.

Generally, top charging is practised, cleaned slack coal being simply run in through the charging holes in the roof of the oven, the charge afterwards being levelled by means of a small ram above the main ram which is introduced through a hole near the top of one of the end doors of the oven. Where weakly caking coals are carbonised, the charge is sometimes "stamped" before being introduced into the In such cases, the crushed coal is compressed by a mechanical stamping machine in a wrought steel box slightly less in size than the oven chamber, and pushed en bloc into the oven as a solid cartridge through the front doorway on the movable bottom of the box which forms the charging peel. At the end of the carbonising period the coke is discharged from the ovens by an electrically driven ram and either quenched by water as it emerges from the ovens or else at a remote quenching station such as will be described later. A good deal of attention has, of recent years, been devoted to dry-cooling of coke with a view to improving its quality and at the same time recovering part of the sensible heat, but wet cooling is still the general practice.

expelled from the coal during the carbonising struction, a short description of the Becker oven

is given.

The Becker Coke Oven.—This oven, which is illustrated in Figs. 4 and 5, is of the vertical flue type and the heating system is arranged so that either lean (blast-furnace or producer) gas or rich (coke-oven) gas may be used. Each oven has a series of vertical flues on either side, under each of which two regenerators are built.

When employing blast-furnace or producer gas for heating, air passes up one of the two regenerators and gas up the other. Both the air and the gas enter their respective regenerators from both sides of the oven at the same time; on leaving the top of the regenerators they enter the bottom of the heating flues so that the air and gas come together with streamline effect and the combustion zone is evenly distributed throughout the length of the vertical flues. The products of combustion are collected from the top of each four or five vertical flues and carried over the top of the oven by means of cross-over flues (Fig. 4), of which there are six to each alternate oven, the intermediate ones not requiring any, as may be seen from the diagram.

From these flues the waste gases are distributed to an equal number of vertical flues on the other side of the oven exactly opposite the flues up which they have travelled. They then pass down these flues and into the regenerators immediately below, through which they pass to the waste gas flues, giving up a portion of their sensible heat to the chequer work in

transit.

At the end of a pre-determined period (from 20 to 30 minutes) the direction of flow of gas and air is automatically reversed, so that the regenerators which have been heated by the waste gases are used to heat the air and fuel gas, while the relatively cool regenerators are heated up. The regenerators are so arranged that those in which gas is being heated are never adjacent to those in which waste gas is flowing, and leakage of fuel into waste gas is therefore impossible.

When rich gas is employed as the heating medium, air passes up all the regenerators and cold coke-oven gas is delivered through a horizontal gas conduit carried in the brickwork between the top of the regenerators and the bottom of the heating flues. From these conduits the gas passes by means of a calibrated nozzle and vertical duct into the bottom of each vertical flue where it mixes with the preheated air coming from the regenerators. The flow of gases from this point is the same

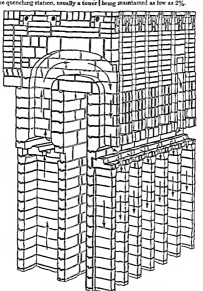
as when blast furnace gas is used.
QUENCHING AND COOLING THE CORE.— A common system of quenching the coke is to spray it with water under pressure while it passes through a cage of pipes on emerging from the oven. The coke then falls on to an inclined coke wharf where it "steams off" and cools and where any hot spots are quenched by

water from a hose.

Where dry cooling is practised, the red hot coke is delivered into a firebrick chamber As an example of modern coke-oven con- through which inert gases (derived from the air

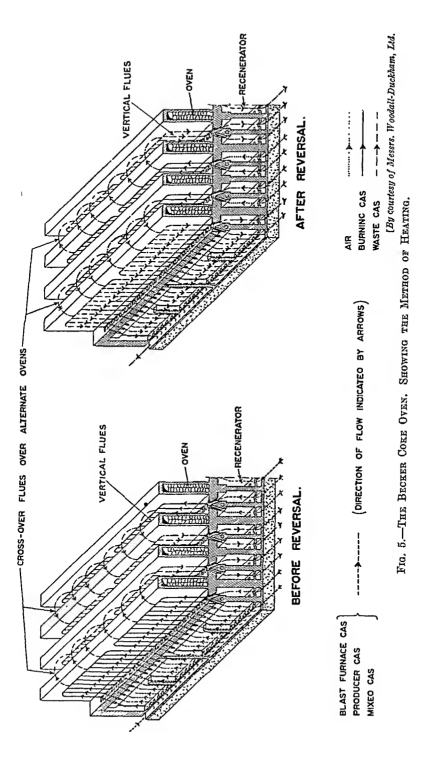
coming into favour; in this process, the coke is while the coke, after draining, is allowed to side discharged into a car which is alowly moved from the car on to the wharf where it steams off across the end of the over, thus spreading and cole finally; by this process uniformity the cole out as a thin layer. The car is rapidly of moisture cortent is secured, the final moisture pushed to the quenching station, usually a tower being maintained as low as 2%.

introduced with the coke) are circulated in [serving also as a chimney to carry away the closed curcuit through the coke and over the large volume of steam produced), where a pre-tubes of a boiler, by which means it is claimed determined quantity of cold water (generally that some 1,000 in of steam (from and at 212°2)] about 3 tons per ton of coke). The practice of remote quenching is now allowed to drain away into a settling pond,



By courtery of Mesers Woodall Ducks am, Ltd. FIG. 4.—THE BECKER COME OVEN. SHOWING THE CROSS OVER FLUE.

Recovery or By-Producers.—Three general systems are in operation, the Indirect, the Direct, obtained in the form of an aqueous solution and the Semi-Direct. The first of these is which must be further teated in order ammonia long or are condensed, while the last traces of tar are aerubited front the gas in a recovery systems are more frequently employed; tar extractor, after which the reidless of the symmonia is removed by sashing the gas with



ammonium sulphate without the condensation of Lquor, or part of the ammonia is directly recovered as sulphate and part AS AD aqueous solution which raust be distilled for recovery of the ammonia.

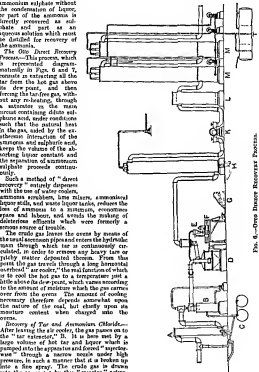
The Otto Direct Recovery Process.-This process, which represented diagrammatically in Figs. 6 and 7. consists in extracting all the tar from the hot gas above its dew point, and then forcing the tar free gas, without any re-heating, through a saturator in the main circuit containing dilute sulphuric acid, under conditions such that the natural heat in the gas, aided by the exothermic interaction of the ammonia and sulphuric acid, keeps the volume of the absorbing liquor constant and the separation of ammonium sulphate proceeds continuously.

Such a method of "direct recovery" entirely dispenses with the use of water coolers,

ammonia scrubbers, lime mixers, ammoniscal liquor stills, and waste liquor tanks, reduces the loss of ammonia to a minimum, economises space and labour, and avoids the making of deleterious effinents which were formerly a serious source of trouble.

The crude gas leaves the ovens by means of the usual ascension pipes and enters the hydraulic main through which tar is continuously circulated, in order to remove any heavy tars or pitchy matter deposited therein. From this point the gas travels through a long horizontal overhead" air cooler," the real function of which air cooler," the real function of which is to cool the hot gas to a temperature just a little above its dew-point, which varies according to the amount of moisture which the gas carries over from the ovens The amount of cooling necessary therefore depends somewhat upon the nature of the coal, but chiefly upon ita moisture content when charged into the ovens.

the "tar extractor," B. It is here met by a large volume of hot tar and liquor which is pumped into the apparatus and forced " injectorwise" through a narrow nozzle under high pressure, in such a manner that it is broken up into a fine apray. The crudo gas is drawn into the apparatus by this "mjector" action, and the sprayed tar and liquor dash against the mechanical spaying of the gas, together with being carbonised, so that the crude gas contains the solvent action of the washing medium, ammonium chlorule, the latter may be removed results in the complete removal of tar, including at the tar parss as a farrly strong aqueous tar fog, from the gas, which passes out of the solution. Indeed, "fixed ammonis" generally extractor in a perfectly clean condition. May, may be removed at this point provided the it should be noted that if a "salty" coal is temperature conditions are such as will allow small tar vesicles contained in the gas. This



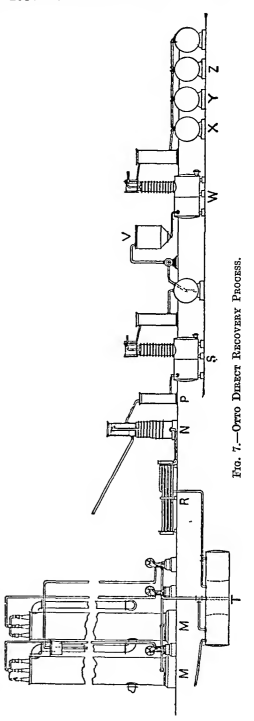
of the condensation of a small amount of liquor in the apparatus. From this the importance of a perfect control of the temperature of the gas, both in entering and leaving the apparatus, will he understood. The tar extracted from the gas overflows into the tar deposit tank, C, which also serves to recover any tars which separate in the long "air cooler," A, already referred to.

Recovery of Ammonia. -- The tar-free gas, which still contains the free ammonia, henzol, and nearly the whole of the naphthalene and water vapour, passes onward, at a temperature (say hetween 65° and 75°C.) so regulated as to he near to (hut not helow) its dew-point, into the large enclosed "saturator," D, containing a hot saturated solution of ammonium sulphate plus an excess of sulphuric acid. In this apparatus, which is lead-lined and cased in a cast-iron framework, the gas is divided into a very large number of small streams, and, under the suction of an exhauster, K, applied at a point further along the system, it is drawn through the seal of the acid liquor, wherehy all the ammonia is removed, and precipitated as sulphate in the coned hottom of the saturator. The exothermic reaction serves to keep the temperature of the gas above the dew-point, so that no condensation of steam occurs in the saturator. The salt is mechanically and continuously ejected from the apparatus by means of compressed air, together with a certain amount of solution, into the cone-shaped receiver, E. The ammonium sulphate crystals settle to the bottom of this receiver, while the hot mother liquor overflows from the top and finds its way hack into the saturator. When this cone-shaped receiver is full of the crystallised sulphate it is passed into a centrifugal dryer, F.

In this machine the sulphate is made thoroughly dry in about three minutes, the mother liquor thus extracted mixing with the above-mentioned mother liquor and with it flows back into the saturator, while the dried sulphate of ammonia is dropped through the bottom of the centrifugal machine, sufficiently dry to be immediately bagged and shipped or put into stock, as shown at G in the

diagram.

Recovery of Naphthalene.—The gas from the saturator, which is now free from hoth ammonia and tar, hut still contains the henzol, moisture, and naphthalene, passes through a lead-lined catch pot (seen as an enlargement in the gas main, whose function is to prevent any acid spray heing carried over from the saturator) and on to the naphthalene extractor and spray cooling tower, H. This tower is quite devoid of any filling, hut at the top there are several jets through which cold water is sprayed at high pressure. The action of this mass of finelydivided cold water beating suddenly on to the hot gas produces a rapid chill, which, together with the concussion brought about hy the water, precipitates and washes out practically over again continuously, as circumstances all the naphthalene, which flows away along permit. This tower also serves to cool the with the water into a catch tank helow. The gases. It is naphthalene is filtered off while the water can be naphthalene, otherwise it would pass on to the



necessary to extract either allowed to flow away or re-cooled and used henzol scrubbers, where it would be dissolved

by the wash oil, which would very soon become reservoir for the circulating benzol wash oil, asturated with naphthalene and rendered useless and the remaining portion being filled with for the purpose of collecting benzol. The naphthalene thus extracted is of a pale yellow colonr, and is readily sold at a price which more than pays the cost of running this part of means the gas is brought into very intimate the plant

Recovery of Benzols.-The gas now being cool. hut still containing the benzel, passes through the gas passes either wholly back to the ovens, the exhauster, K, and a second water cooler, L, or in the case of regenerative ovens, about which takes out the last traces of naphthalene, one-half of it returns to the ovens, and the and thence to the benzel scrubbers, MM, which other half is used either for firing boilers, vary in number according to the size of the town lighting, operating gas engines, or other plant. These scrubbers are from about 50 ft. purpose.
tr 70 ft. high, the lower portion forming a According to E. Bury (Trans. Inst. Min. Eng.

wooden grids. Benzol wash oil is constantly circulated downward over these grids, the gas having to pass upward through them. By this contact with the oil, with the resulting absorption of the benzol. After leaving these scrubbers

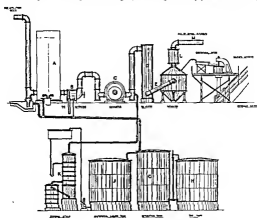


Fig. 8.—The Kotpers Semi-Direct Recovery Process.

1911, p. 537), the following temperatures were passes downward over a series of trays. Dr. observed in a plant at New Brancepeth Colliery (Co. Durham):

Hydraulie main	225	
Before tar extraction	86	
After tar extraction .	58	
In anturator	67	
After saturator	58	
After exhauster	68	

Distillation and Rectification of Benzole.-The " wash oil" containing the benzols in solution is sent to the continuous crude benzol still, passed through the refrigerator, R, whence, N (Fig. 7), where, after being pre-heated by the after being cooled to the atmospheric tempera-

ateam at about 100 lb. pressure, reduced, however, at the still to 10 lb. pressure, is delivered into the hottom of the apparatus, and passing upwards bubbles through the wash oil, liberating the benzol hydrocarbons which it carnes over into the condenser, P. Hero the ateam and benzols are condensed, and the condensed water and oil separated. The crude benzol then either passes into a storage tank, or (if its rectification is to be carried out) into the primary rectifying still," S. The debenzolised " wash oil" leaving the crude benzol still is hot vapours coming away from the still, it ture, it is sent into the "oil storage tank," T,

and from there it is used again in the henzol!

The erude henzol is first of all fractionally distilled in the primary rectifying still, S, 3,000 to 5,000 gallons capacity and intermittent in its working, in which it is heated under reduced pressure hy means of steam coils, and, with the aid of a long dephlegmating column, it is separated into:

(1) Crude 90% Benzol.
 (2) Crude 90% Toluol.
 (3) Crude 90% Xylol.
 (4) Crude 90% Solvent Naphtha.

Each of these fractions is separately condensed and pumped into a large vessel, V, termed the agitator, where it is successively treated with (a) pure strong sulphuric acid, and (b) a solution of caustic soda, in order to remove all resinous, basic, or acidic substances. Each of the A, where the tar and water vapours are con-

"washed" fractions is finally refractionated under reduced pressure in the "secondary rectifying still," W, and the re-distilled resulting fractions, after condensation, are sent forward into their respective storage tanks, XYZ, ready for transport.

Other firms (e.g., the Simon-Carvés, the Collin, and the Simplex Companies) have also adopted efficient "direct recovery" processes, on much the same lines as the Otto process just described, the chief difference between them heing in the precise mechanical arrangement for the removal of tar and tar fog from the hot gas before it enters the saturator where the ammonia is absorbed.

The Koppers' Semi-Direct Recovery Process.— In this process (illustrated in Fig. 8), the hot gases from the ovens are first cooled approximately to air-temperature in the primary coolers,

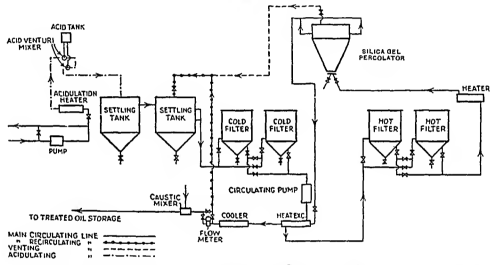


Fig. 9.—Diagram of Plant for the Refining of Benzol by Means of Silica Gel.

exhauster, C, through the tar-extractor, B, where the last traces of tar fog are mechanically removed. The tar-free gases are next heated to ahout 50° to 60°C., in the re-heater D, by means of exhaust steam obtained from the engines used in the recovery plant, and are then brought into contact with sulphuric acid in the saturator, F, where the ammonia is removed as ammonium sulphate.

The tar and ammonia liquor which are extracted in the primary cooler and tar extractor are collected and delivered into the decantation tank, G, where separation takes place, the tar and liquor heing separately run into the storage tanks H and J. The liquor is continuously pumped into the ammonia still, K, where the free and fixed ammonia are driven off hy steam and lime. The vapours from the still are returned into the gas-stream and pass with it into the saturator.

After leaving the saturator, the gases pass through the acid-separator, L, where any acid

densed. The gases are then drawn by the removed, and thence are conducted by the main, M, to the henzol-recovery plant which operates on the same general lines as that already described.

As an alternative to the oil-washing process the recovery and refining of benzol hy solid adsorbents (silica gel or activated carbon) has been developed. For the use of activated charcoal for this purpose, see end of article. E. C. Williams (Proc. Chem. Eng. Group, S.C.I. 1924, 6, [A], 95-114) discussed the principles of the recovery of benzol hy silica gel and described an experimental plant designed by the Silica Gel Corporation. This process does not yet appear to be operating on the large scale, the main difficulty being the serious wastage of the gel; silica gel, however, is employed in the refining of henzol, and this process is dealt with helow.

Benzol Refining by Silica Gel.—The silica gel process for benzol refining is primarily catalytic, polymerisation of the gum-forming impurities being initiated hy acid treatment, the gums are spray carried forward from the saturator is then deposited on the surface of silica gel, where

they act as catalysts to polymerise the remain-1 ing unsaturated hydrocarbons. The crude The similarity in composition of the gas benzolis pumped through a tubular ateam heater of the conduction of the conduction of the conduction heater in Fig. 9, p 267. Thence at type already described or in a vertical grant the raise its temperature to about 40°C. Thence at type already described or in a vertical grant of the conduction of flows through two Venturs meters; while passing (as illustrated in Table II) is such that the fields the first, 0 1% of concentrated sulphuric acid is of utility of coal- and coke oven gases are likely added to dry the benzol, and at the second to be the same. meter, 0.2% to initiate polymerisation. After passing through two settling tanks where a Table II.—Compositions of Core-Oven and amall amount of sludge is drawn off, the acidulated benzol passes through a coke filter, and is then pumped to a heat exchanger, where it is heated by hot benzol from the sales gel perco-lator, and is afterwards filtered while hot through the "hot coke filter." After being heated to about 150°C. (under pressure aufficient to maintain it in liquid phase), the benzol passes upwards through the conical sibes gel percolator where polymerisation and removal of gumming constituents are completed, it passes through the heat exchanger counter-current to the benzol from the circulating pump, through a cooler, and flow meter, and finally, after treat ment with caustic sods to remove acidity, to storage. After the silica gel percolator has been used to treat 1,500 times its own weight of henzol, it is removed for reactivation, and replaced by a freahly reactivated

chargo. In dealing with the subject of by-product recovery, mention should be made of the fact that the rapid development of the synthetic ammonia industry has led to a fall in the price of ammonium sulphate which has rendered ammonia recovery from coke oven gas so much less profitable than formerly that in some cases it has been abandoned. In Table I the pro-duction of ammonium sulphato and ammonia liquor at coke oven plants in the United Kingdom in 1924 and 1930 are compared.

The quality of ammonium aulphate now marketed is higher than that before the War, the salt being neutralised and dried before being placed in atorage. The apecification of the British Sulphate of Ammonia Federation demands that the salt should fulfil the following requirements: Nitrogen content to be not less Durham 25652025055095 than 20.60%; free scudity (calculated as H₃SO₄) not greater than 0 025%; and mossture S. Yorks 2 4 5 8 2 0 26 4 54 6 9 1 not over 0 20% (±0 05%).

TABLE I .-- PRODUCTION OF AMMONIUM SUL-OVEN PLANTS IN THE UNITED KINGDOM.

	1924.	1930.
Total coal carbonised in by- product ovens, milion		
tons	17 23	1651
Ammonium aulphate pro- duced, tons	186,200	173,100
Sulphate produced per ton of coal, lb	242	23 5
Ammonia liquor produced,	12,700	5,800

THE UTILISATION OF COKE-OVEN GAS.

VERTICAL-RETORT GASES BEFORE BENZOL

MECOVERY.		
	Coke oven gss. Mixture of 8 Norkshire and Derbyshire eonis carbon- ised in Becker ovens	Vertical retor gas Mitchel Main gas nui carbonised in Glover-West vertical retort
Percentage: CO: CO: Co: CoHm CH; H; O: N: Alortice Value	2 2 6 8 3 0 28 2 52 6 0 8 6 4	0 8 7 4 2 2 29 05 52 0 0 9 7 95
(B Th.U.) per cu ft.(grossat 15°C) and 760 mm.	553 4	554 0

Debenzolised Gas -The amount of benzol vapours included among the unsaturated hydrocarbons, CnHm, in the above table would be approximately 1% in each case, and the effect of removing them would be to reduce the caloratic value of the gas by from 5 to 10%. The average percentage composition of the "debenzolised" gas from typical Durham and South Yorkshire coking coals is somewhat as follows:

Catorific values (B'Ih U) ខីខភិ ភិ ភ 15°C and 760 mm 1481 9 gross 1425 5 net (493 6 gross | 434 2 net

Amount of Surplus Gas .- Taking the country as a whole, and assuming that the whole of the PHATE AND AMMONIA LIQUOR AT CORE 17-4 million tons of coal carbonised in 1935 for metallurgical coke was dealt with in regenerative ovens, the net calorific value represented by the surplus gas (at 5,500 cu. ft per ton) available for outside purposes might have been about 47,300 therms per hour, day and night, or aufficient to generate about 490,000 B H.P. continuously in internal combustion engines.

The most effective utilisation of such a hugo aurphus of energy is a matter of the highest importance to the nation, and there is no doubt that a good deal of the coke oven gas available from existing plants is not being used to the best advantage.

Table III, taken from the reports of the Secretary of Mines, shows the make of cokeoven gas and how it was actually used during purposes by the oven owners, for steam raising | 2½% unaccounted for.

the period 1932 to 1935. In 1935 the average or on steel works where the ovens and steel gas production was 10,870 cu. ft. per ton of coal, of which 59.5% was used for heating the 11% was sold to gas companies and 3.5% to ovens, while a further 23.5% was used for other other undertakings, leaving a balance of about

TABLE III.—By-PRODUCT COKE-OVEN GAS PRODUCTION AND DISPOSAL.

Year.	1932.	1933.	1934.	1935.	Cu. ft. per ton of coal, 1935.
		Million	cu. ft.		
Total produced	124,230	137,492	177,070	185,558	10,870
Used by Oven Owners. For heating ovens For other purposes	89,092 {	81,269 31,790	106,519 40,146	110,236 43,731	6,460 2,560
Sold to Gas undertakings Other undertakings	15,480 15,994	15,941 6,240	18,052 8,276	20,474 6,327	1,200 370
Balance	3,732	2,251	4,078	4,791	280

In normal times by far the greater part of the coke manufactured outside gasworks is for metallurgical purposes, and chiefly for iron-smelting. In cases where blast furnaces are taken place in European countries, and particular to the countries of the countries attached to steel-works, the utmost fuel economy may be achieved by concentrating coke ovens, blast furnaces, steel-furnaces, and rolling mills as a self-contained plant on one site so that combined surpluses of coke-oven and blast-furnace gases may be utilised, thus making it possible to produce finished steel sections from the ore with no more fuel than the coal necessary to furnish the coke required in the blast furnace. In such cases there is usually no surplus cokeoven gas available for outside purposes except at week-ends, when the steel-works and rolling mills are shut down.

Such considerations do not, however, apply to independent blast furnace plants producing foundry iron (e.g. in the Midlands) or to highgrade steel-works (e.g. the Sheffield area) unattached to blast furnaces. In such areas the coke oven plants are usually situated at the collieries, and large surpluses of gas are available for outside purposes. Thus it has been estimated that in 1928 within the area comprising South and West Yorkshire, Lancashire, Derbyshire, Lincolnshire, Cheshire, Notts, Staffordshire, Leicester and Warwickshire, there were available for "outside" purposes about 10,000 million cu. ft. of coke-oven gas of an average calorific value 500 B.Th.U. per cu. ft., of which about 4,200 million cu. ft. were sold to gas undertakings, and another 2,150 million cu. ft. to other undertakings. Moreover, with new coking plants then contemplated, it was considered that the total available surplus gas within the area might soon amount to as much as about 22,500 million cu. ft. per annum. Such a prospect, being subject to fluctuations in the activity of the iron and steel industry, is therefore to some extent doubtful, which is a serious drawback from the point of view of network either in Lancashire or West York-

ticularly Germany, in regard to the pooling and distribution over large areas of such surplus coke-oven gas; and the question arises whether or not some such schemes would be equally practicable in certain areas of Great Britain, seeing that nowadays there would be no technical difficulties to overcome. Thus, for example, at the end of 1929 in Germany the Ruhr Gas A.-G. (established in 1926) owned 286 miles of welded steel mains, with a further 330 miles under construction or contemplated, and in 1930 distributed some 21,000 million cu. ft. of coke-oven gas; moreover, in 1929, about one-third of the total gas consumed (i.e. from gasworks and coke ovens) in Germany was derived from the Ruhr coke ovens. The gas is guaranteed free from tar and H₂S, the maximum permissible content of NH₃ being fixed at 0.87 g. and of naphthalene at 4.37 g. per 100 cu. ft. The price delivered to gas companies is usually based on a sliding scale according to the price of coal (e.g. to Hanover at the equivalent of 11.6d. per 1,000 cu. ft. up to 1,765 million cu. ft. per annum, and at 10.6d. per 1,000 cu. ft. for consumption exceeding 2,472 million cu. ft. per annum, when the price of coking coal is equivalent to 18s. 9d. per ton), but to industrial establishments the price is equivalent to 10d. per 1,000 cu. ft.

In 1930 a Departmental Committee of the Board of Trade was appointed to inquire into the technical and economic aspects of an area gas supply comprising the afore-mentioned counties in England, and, after hearing much evidence, reported that while circumstances would not warrant the construction of elaborate and expensive systems of trunk mains over the whole area, and are not yet favourable for such

shire, tha iron and steel industry around electric energy which is distributed for public Sheffield and Rotherham offers a good market consumption over a wide area, as in the Northfor all the surplus coke-oven gas likely to be East Coast Power Scheme available in the near future if it could be supplied at an averago price of 2d per therm (-say 10d. per 1,000 cu. ft.). Accordingly they recom-mended the adoption of a scheme for such restricted area embedying the following conditions (inter alia); (i) To collect gas from 19 coke oven plants and to take a maximum volume of 80 million cu. ft of gas per day, tha flow of gas to be uniform throughout the 24 hours of the day, (a) the gas tu be purified at the coke ovens before delivery into the network; (iii) the compressor plants to be in-stalled at the coke overs and their cost and the cost of operation to be met by the coke-oven owners, such costs being covered by them in their price of gas; (1v) the pressures to be used to vary from 2 7 lb to 8 2 lb per sq in , when the mains are estrying up to 40 million eu ft. per day, and from 49 lb to 22 lb, when they sro carrying up to 80 million cu. ft, per day; (v) the length of the mains to be approximately 73} miles, and to be of steel of medium weight varying from 6 in, to 30 in in diameter with hessian cloth and with bitumen coating, and facture of synthetic ammonia at Ostend, wem. laid in roads, receiving 3 ft. cover, at an estimated capital cost of approximately £537,000, (vi) the cost of distribution (exeluding compression) to vary from 1 33d. per 1,000 eu ft. (0 25d, per therm) with a voluma of 40 million eu. ft. per day to 066d. per 1,000 cu. ft (0 13d. per therm) with a volume

of 80 million cu ft per day It was further considered that the con-ditions of gas supply in Germany (where town's gasworks production is less than half that in Great Britain, and the consumption per head of population about 1,800 cu. ft. per annum as compared with over 6,000 cu. ft. in Great Britain) "aco so dissimilar to these in the country that it does not follow that longdistanca transmission of gas can be introduced into Great Britain on the German model."

Among other directions in which the surplus gases can be ntilised to good advantage may be mentioned:

For Steam Raising and Power Purposes .-By burning the gases in the tubes of a Bonecourt Boiler, on the principle of " surface combustion," it is possible to tronsform 92% of the net heat of combustion into the energy of atesm which can be either (a) converted into mechanical power by means of a steam turbino, or (b) used for evaporative purposes in connection with the definition of the state of net calonfic value of the gas to B.H.P. at the importance in connection with motor fuel engine). And by suitable combinations of production. In this country some 33 million either (c) believe, turbunes, and dynamos, or imm of coal are carbenized annually by the gas (d) gas engines and dynamos, electrical energy and coke-oven industries, and in 1934 tho ean be generated at s very low cost. Many annual total production of refined benzol collieries now generate from their aurplus coke- from all sources had reached just uver 32 million oven gases (or waste heat) sufficient power tu gallons. This production could be more than meet all the requirements of the mine. Others, sgain, sell their surplus gas (or waste heat) sgam, sen their surplus gas for waste seat) second Internation to a power company, which converts it into Coal, Pittsburgh (1928).

If no power company, or group of power users, is at hand to purchase the surplus gas (or waste heat), it might with great advantage be utilised m an adjacent chemical factory. For thero is no reason why the crude ters, benzols, and naphthalens produced on the coking plant should not be refined and transformed into other more valuable products in a chemical factory on the spot, the heat and power for which would be wholly derived from the surplus gases. Or, alternatively, any chemical operations requiring cheap electric current might very well be carried out in factories erected in closo proximity to

hy-product coking plants. As a Raw Material in the Chemical Industry .-Coke uven gas is an important source of the hydrogen used in the synthetic ammonia industry. Thus Pollitt 1 stated that 15 8% of the total amount of hydrogen used throughout the world for ammonia synthesis in 1930-31 was derived from coke oven gas. According to Pallemaerts, tha relative costs of preparing hydrogen at a pioneer installation for the manu-(1) From water-gas (from coka at 10 6d. per ton) 8 7d. per 1,000 eu. ft , and (u) from cokeoven gas (at 5 45d per 1,000 cu ft) 9 27d. per 1.000 eu ft.

The residual gas, after removal of hydrogen, has a calorific value some 25% higher than that of the coke-oven gas, and could be delivered into the mains for industrial purposes. On the other hand, during the preparation of hydrogen from the gas, three fractions are obtained, (1) a fraction containing about 18% of carbon monoxide and 73% of nitrogen, which is used on the Continent for the synthesis of methanol,
(u) a fraction containing about 75% of mothane,
and having a calorific value of about 815
B.Th U. per cu. ft., which would be suitable
as a raw material for synthetic reactions, and (m) a fraction containing approximately equal parts of ethylene, ethano, and methano, together with a little propylena and carbon menoxide, which is used at Bethune for production of ethyl and propyl alcohols

Laterature -W. A. Bono and G. W. Himus, "Coal, Its Constitution and Uses," Chap. XXII, 1936; and J. Roberts and A. Jenkner, "International Cost Carbonisation,

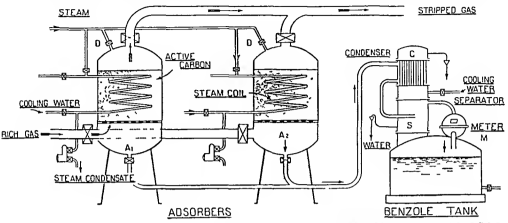
" 1934. W. A. B. and G. W. H.

BENZOL RECOVERY BY ACTIVE

World Power Conference, Berlin (1930)
 Second International Conference on Bituminous

present at a partial pressure of 7 to 8 mm. | processes on a large scale. Prior to 1929 the

doubled if all the coal gas made were submitted | Formerly the recovery of this benzol was imto treatment. In the carbonisation of 1 ton of coal about 3 gallons of benzol are formed. Part of the benzol is found in the tar, but the illuminating standards in the gas industry have bulk of it accompanies the gas in which it is rendered possible the practice of extraction

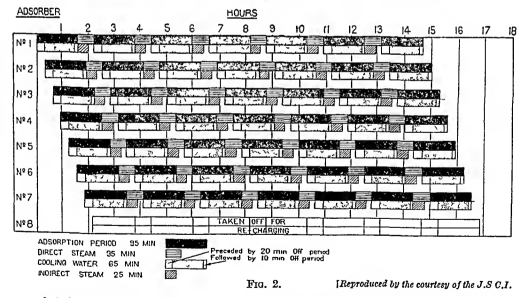


Frg. 1.

[Reproduced by the courtesy of the J S.C.I.

only commercial method in this country was | that of absorption by heavy oil, which is still most widely used. The active charcoal process was originated by the Bayer Company (G.P. 310092, 1916; B.P. 156543, 1916; A. Engelhardt, Gas-u. Wasserfach, 1921, 64, 205; 1922, 65, 473) and is fairly well known on the Continent. A disability of the process as first (H. Hollings and S. Hay, J.S.C.I. 1934, 53, introduced was the short working life of the 143). The activo charcoal process has certain

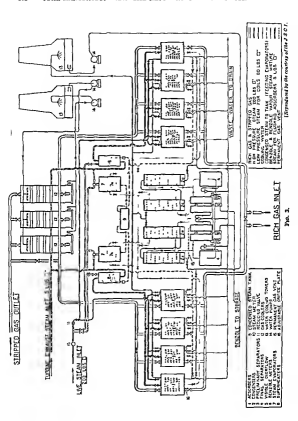
adsorbent, but this difficulty has now been largely overcome (H. Hollings, S. Pexton and R. Chaplin, Trans. Inst. Chem. Eng. 1929, 7, 102; Gas J. 1929, 188, 715). In 1929 the first commercial unit was installed by the Gas, Light & Coke Company at Harrow, and a few



technical advantages over the oil process, and is | treated must be free from tar fog, hydrogen

bledy to be more widely adopted in the future.

Description of the Process.—Chemically activated charcoal of high volume activity and low retentivity is used in the form of short rods or pellets 4 to 5 mm. in diameter. The gas to be Gas passes upward through adsorber Al,

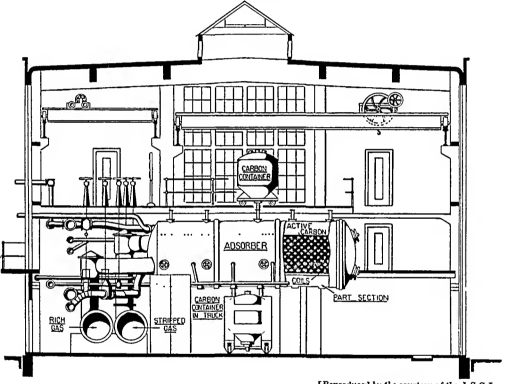


containing active charcoal, which retains henzol and certain other constituents of the which retains! gas (e.g. unsaturated hydrocarbons and carbon disulphide), cold water heing circulated through the coils at the same time. When the charcoal in Al is saturated the gas is diverted to adsorber A2, while A1 is regenerated. For this purpose the cooling water is shut off and steam is passed through the coils in order to heat the charcoal. At the same time direct steam is passed downwards through the charcoal hed to remove the adsorbed henzol. The mixture of steam and henzol vapour leaving the adsorher passes to the condenser C, and the condensed liquids are separated in the gravity separator S, from which the henzol flows through the meter M to storage tanks, while the water runs to waste. When can be operated as a single unit or independently. steaming is completed gas is again admitted to the hot adsorber which removes excess moisture contains four adsorbers. The adsorbers (Fig. 4)

and cools the charcoal rapidly. After a short time cooling water is again circulated through the coils and adsorption carried to completion. The cycle of operations is thus reperted. The size and number of adsorbers required are determined by the volume of gas to he treated in unit time, hut not less than two adsorbers can he used if the adsorption process is to be

continuous (see time schedule, Fig. 2 on p. 271).

THE BECKTON PLANT.—This installation deals with gas from horizontal retorts and coke ovens and has a potential output of 20,000 gallons of benzol per day, recovering 2.9 gallons from 10,000 cu. ft. of gas with an efficiency of 92–95%. The layout of the plant is shown in Fig. 3. It is arranged in two halves which Each half has its individual gas inlet main and



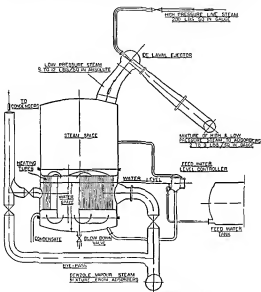
[Reproduced by the courtesy of the J.S.C.I.

Fig. 4.

are cylindrical vessels, 9 ft. in diameter and 27 ft. in length, and hold 7 tons (700 cu. ft.) of active charcoal, which is supported on hinged grids. The pipe coils used for heating and cooling. The charcoal is charged into the adsorbers from closed containers which are run into position on overhead trucks. For discharging the charcoal the grids are lowered at the centre by means of special gear and the spent material received into similar containers placed underneath. Communication between the two levels is effected hy a hydraulic lift. Overhead cranes are Vol. III.-18

provided for assembling and dismantling parts of the plant. Each half of the plant is provided with a steam regenerator, a set of tubular condensers, a set of preliminary separators, a final separator, and a henzol meter. The stripped gas is cooled by direct contact with water in three gas coolers which condense the water removed from the hot charcoal after steaming. Cooling water for all purposes is supplied by electric pumps, but the water circulated through the adsorber coils is contained in a closed system charged with de-aerated, condensed steam to minimise corrosion.

OPERATION OF PLANT.—There is considerable either in reserve or being recharged. Gas is latitude in the rate of passing gas which may passed until the benzol "hreakpount" as vary from 300,000 to one milion cu ft, per reached as indicated by fist flame burners fed hour per adsorber. The number of adsorbers at from the outlet stream. The stripped gas at work is governed by the amount of gas being first burns with a practically non-luminous made and by the age, i.e. activity, of the flame, but as little as 0.04 gallon of benzol charced in two. At full capacity five are made per 10,000 cu. ft. of filtered gas shows Ma a gas, two under steam, and the remaining one luminous fringe in the flame. In steaming the



[Reproduced by the courtery of the J.S. C 1. Fig. 5.

direct and indirect ateam supplies are turned on cycle cooling water is not used for the first simultaneously. The adsorbed gates first evolved 20 minutes, so that the charcoal may be contain bergol and are returned to the subst sufficiently direct by the rich gas. consam sested and are remease to me much suggested upon 10 to the gasstream. After 5 or 6 minutes, at a temperature of 5 sfCo. bennol commences to be evolved. Steaming is continued 30 to 40 minutes, according to the age of the charcoal, which is finally steam from high pressure mains and partly of ing to the age of the charcoal, which is finally steam from the team requirement of 102, 50, 10.

Taked to a temperature of 12 2°C, and contains

Cheap rhant steam may replace a considerable

15% moisture. In the subsequent adsorption part of the high pressure steam when two

from high pressure mains after reduction to 80 lb. pressure, and de-superheating. Requirements and distribution of process steam are shown in the following table.

	Steaming one adsorber.	Steaming two adsorb- ers at once.
Indirect, to coils Direct steam : (a) High pressure	lb. 1,000	2,000
steam (b) Regenerator steam (c) Exhaust steam .	6,000 4,500 —	6,000 5,000 10,000
	11,500	23,000

About 6% only of the total heat requirements of the process are accounted for by the latent and sensible heats of the benzol. The rest is used in raising the temperature of the adsorber and contents. The average overall steam consumption is about 34 lb. pcr gallon of benzol.

THE STEAM REGENERATORS .- These are an important feature of the plant and supply 32% of the heat contained in the distillate leaving the adsorbers. A regenerator is shown diagrammatically in Fig. 5. Distillation vapours from the adsorbers pass between a nest of tubes between two header plates before entering the condensers. A De Laval steam ejector in communication with the top of the chamber produces a partial vacuum and lowers the boiling-point of the water in the tubes so that heat is transferred from the vapours to the steam produced which is entrained by the high pressure steam in the ejector. The feed water consists of condensate from the adsorber coils augmented by softened water from outside sources and is maintained at constant level by a float valve. The two regenerators supply between them 4,500 lb. of steam per cycle.

PRACTICAL ASPECTS OF THE PROCESS.—The adsorptive capacity of the charcoal varies but slightly with changes in benzol concentration in the gas, e.g. an increase from 2.13 to 2.85 gallons per 10,000 cu, ft. only increases the capacity from 32 to 33% by weight of adsorbent. Since the "slip" of benzol is unaltered the recovery efficiency is also increased. The recovery efficiency is also increased. The adsorptive capacity is considerably lowered by the presence of moisture in the charcoal or in the gas. It is therefore an advantage to have the gas as dry as possible.

The efficiency of recovery of benzol increases by about 1.2% per degree C. fall in temperature; hence one reason for cooling the charcoal during adsorption. The heat of adsorption of benzol (340 B.Th.U. per lb.) is sufficient to raise the charcoal temperature to 140°C., but this is effluent gas, etc., that the adsorption tempera-

adsorbers are steamed simultaneously, as the the temperature from 120°C. to 60°C. in 20 table shows. Indirect or coil steam is taken minutes, and is of great importance in prolonging the working life of the adsorbent.

The last traces of benzol are difficult to remove from the charcoal and practically desorption is at an end when the ratio of direct steam to recovered benzol is 10:1, i.e. after about 20 minutes. At this point, however, the charcoal temperature is too low for efficient drying, so that steaming is continued for a further 10 to 20 minutes (the newer, i.e. the more active, the adsorbent, the greater is the amount of steam required).

DEPRECIATION OF THE ADSORBENT .- Hydrocarbons of high molecular weight and certain unsaturated hydrocarbons adsorbed from the gas are preferentially retained by the charcoal after steaming. These accumulate in charcoal and gradually impair its activity towards benzol. The unsaturated hydrocarbons are particularly troublesome in this respect because when the charcoal is heated they polymerise into gummy substances for which the adsorbent has a high capacity, and which can only be removed by oxidation at high temperatures. The rapid deterioration of the adsorbent proved to be a serious disability of the active charcoal process which was at first uneconomic through this cause (H. Hollings, S. Pexton and R. Chaplin, *l.c.*). Operative procedure has now been so much improved that the adsorbent can be used from 1,500 to 1,900 times, and made to recover about 175 times its own weight of benzol before needing replacement. The following practices are mainly responsible for this improvement:

(i) the passage of steam countercurrent to gas;

(ii) the simultaneous use of direct and indirect steam at high rate, which limits the heating period;

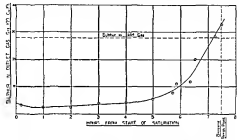
(iii) the adjustment of a critical moisture content in the charcoal after steaming so that it cools rapidly in contact with rich

In this connection the employment of charcoal of low retentivity, i.e. of low capacity under the desorption conditions of the process, is obviously important.

Spent charcoal contains about 33% of fouling material, but may contain as much as 45%. A more or less complete regeneration may be brought about by the action of air or of steam with air at 500-600°C. (B.P. 313154 and 338500, 1929). In these processes the fouling matter is carbonised and burnt away but a certain loss of adsorbent is inevitable. recent process based on hydrogenation of the fouling material under pressure, and which entails no loss of adsorbent, is described under B.P. 374597, 1932.

SULPHUR REMOVAL.—In adsorbing benzol active charcoal also removes sulphur compounds dissipated at such a rate by the coil water, from the gas. Practically all the carbon disulphide and thiophen are removed in the ture remains in the region of 20°C. The most early stages, but as the benzol breakpoint is effective cooling is provided by evaporation of approached some carbon disulphide and residual moisture in the adsorbent which lowers other volatile sulphur compounds are dis-

placed into the filtered gas. Fig. 6 shows in a 75-80%, which compares favourably with parcel amonte the variation of total subject of the processes expedited by extent content of the subject of the subject of the subject of the processes expedited by existed for subject or the subject of the subject o



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(i) high recovery efficiency; (u) low steam, water, and power consump-

tion: (iii) purification of the gas from sulphur.

From the economic point of view there seems little to choose between the active charcoal and oil processes (v. Adsorption, Vol. I. p. 150a)

COLAMINE v. CHOLINE.

COLCHICEINE v. Colchicum.

COLCHICINE e. COLCRICUM.

COLCHICINIC ACID v. Colchicum.

COLCHICUM and COLCHICINE. Colchicum autumnole Lina. (Fam. Laliacen), Meadow

saffron (Fr. Colchique, Ger. Herbstzestlose); the plant is found in meadows and pastures throughout Europe, where its crocus like flowers, usually of a lilac hue, occur in September and October.

The Brussels conference agreed that the seeds only should be used; in the "British Pharmaco-" 1932, the seeds and corms are official; in the "United States Pharmacopæia," 1936, the seeds and the alkalold colchicine are official. The alkaloidal content of the plant varies greatly, mostly from 0-2 to 0'8% in the aceds ("British crystalline powder melting between 135" and Pharmacoporia"; 1932, not less than 03%; 145° is obtained. The aalicylate is then usually "United States Pharmacoporia," 1936, not less prepared. then 0.45%), from 0.1 to 0.5 in the corns ("British Pharmacopeia," 1932, not less than Pharmacopeia," 1932, p. 131; and "United 0.25%), Colchience occurs to a sight extent [Sates Pharmacopeia," 1936, p. 127. Further also in the flowers.

same genua and also in the large tubers of Grier (shid. 87).

consumption of ateam per unit of benral (Glorious superios (Fam. Lalances), common in India and Ceylon (0.99) of "gloriousne,"
Points in favour of the active charcoal process (Clever, Green and Tutin, J.Cs. 1915, 107, 399). Klewn and Pollary (A. 1931, 778), found colchicine in considerable quantities in the following Liliacem: Bulbocodium, Tofieldia, Veratrum anthericum, Hemerocallis, Ornithogalum, Tulipa; cf. also Albo (J.C.S. 1901, 80, II, 679). Fourment and Roques (Amer. Chem. Abetr. 1927, 21, 2166) isolated colchicine from Merendera Bulbocodium (Fam. Liliacem)

Extraction of Colchicine .- The alkaloid occurs as free base in the plants; all preparations of colchicine are based on its removal from acid ur neutral solution by CHCl₃. Chemnitius (J. pr. Chem. 1928, [11], 118, 29) proceeds as follows: Ground seeds are exhausted with hot 80% EtOH and the solvent evaporated. Fat and resum are removed by addition of molten paraffin wax to the still warm concentrated solution, which when solidified can be removed mechanically. Water is added to precipitate further amounts of tarry material. From the aqueous solution the hase is then shaken into CHCI, this solvent evaporated and last traces removed by distillation with absolute EtOH.

The chloroform-free alcoholic solution is filtered and the alkaloid precipitated with four times its volume of Et.O. when a alightly yellow semi-

references, see Self and Corfield (B. 1932, 1135). Colchicine is present in other species of the Hooper and King (Pharm. J. 1923, 111, 104),

with platinirhodanide, see Klein and Pollauf (l.c. 1931), colour reactions of colchicine, see Ekkert (A. 1926, 533; ibid. 1929, 86), estimation of colchicine with phosphotungstic acid, see Davies (Amer. Chem. Abstr. 1921, 15, 3894).

Colchicine (I), $C_{22}H_{25}O_6N$, $[a]_D^{16.5}$ -120·6° -429° (in H_2O), m.p. CHCl₃) and 155–157°, soft pale yellow needles CH3 CO2Et (Clewer, Green and Tutin, l.c. 1915). The alkaloid is usually seen as a yellow anhydrous varnish, m.p. 142°-147°; Zeisel first crystallised it from CHCl₃ with two molecules of CHCl₃ of crystallisation; according to Merck (Pharm. Zentr. 1916, 57, 619) there are two such compounds, B.CHCl3 and B2.CHCl3. The crystalline colchicine is mostly the latter compound in which 14-16% CHCI3 is retained tenaciously, and is only given off slowly at 100°. According to Merck, pure (I), dissolved in 3 parts of H2O, slowly crystallises as the bydrate B_2 ·3 H_2O .

(I) dissolves slowly but abundantly in H₂O, readily in dilute EtOH, CHCl₃, and hot benzene, less readily in bot H₂O (12%) or absolute EtOH, hardly at all in Et.O.

Being an acid amide, (I) is a very feeble base and one of the few alkaloids which can be extracted from acid solution by CHCl₃. No salts anhydrous m.p. 172°; it gives a green

For microchemical identification of colchicine are formed except an aurichloride, m.p. 209°. Dilute mineral acids and alkalis colour colsolutions intense yellow. cbicine (d 1.4) colours the solid alkaloid a deep violet, changing to yellow and finally to green (charphosphomolybdic Tannic acid, acteristic). acid, potassium triiodide, and bismutb potassium iodide are the most sensitive precipitants. For dissociation of (I), see Kolthoff (Biochem. Z. 1925, 162, 348); for absorption-spectrum of (I), see Purvis (J.C.S. 1927, 2715).

Constitution of Colchicine.—Early work on the constitution of (I) is due to Zeisel and co-workers (Monatsb. 1883, 4, 162; 1886, 7, 557; 1888, 9, 865; 1913, 34, 1327, 1339), but only the brilliant research of Windaus and co-workers gave definite information (J.C.S. 1911, 100, Ī, 904; 1915, **108**, I, 708; 1924, **126**, I, 72; Annalen, 1924, 439, 59; Ber. 7924, 57, [B],

1871, 1875).

(1) contains a N-acetyl and four MeO-groups; if it is beated for a short time with 0.5% HCl or weak alkali, colchiceine (II) is formed with loss of one MeO-group. Contrary to (I), which is a practically neutral substance, (II),

C21H23O6N, H2O,

an enolic form of a β-keto aldehyda (keto hydroxy methylene grouping, tautomeric with

an o hydroxy-aldchyde).

By methylation (11) is transformed back to colchicine and N-methyl colchicioe. By the action of hypotodite the enolic aldehyde group in (II) is replaced by iodine, a transformation in (11) is repeated by former at transmission characteristic of o and p bydroxy aldehydes, the resulting compound is called N acetyl-iodocolchnol, CostH.co.N.1 (11).

If (1) is heated with 15% HCl the N acetyl group is removed as well as one MeO. group.

and trimethylcolchicinic acid, CieHarOaN (IV), results. The latter compound yields salts with alkalis and acids. By tresting (IV) with hydroiodic acid, the remaining three MeO groups are

removed and colchicinic acid results.

By oxidation of (1), (11), (111), or (1V) with permanganate 3 4.5 trimethoxyphthalic (from ring 1) and succinic acids are obtained. By potash fusion and aubsequent permanganate oxidation terephthalic and trimellitie (V) (1:2.4-benzenetricarboxylic) acids are obtained, indicating the presence of a second six membered

ring (ring 2)

N-acetyl iodocolchinol (see above) has a phenohe OH group and can easily be methyl ated. If this methyl ether is oxidised first with nitric acid and then with permanganate, rings 1 and 2 are destroyed and todo methozy phthalic acid (VI) is isolated from the exidation mixture This latter acid originates from the third aix-

membered ring present in (I).

The methylether of (III) (see above) can be reduced (elimination of iodine) and hydrolysed (elimination of acetyl group) to colchinol-methylether, C₁₉H₂₃O₄N; this compound is then submitted to a Hofmann degradation, eliminates NH, and generates double bond in the central ring, forming 2:3:4:6 tetramethoxy · 9 · methylphenanthrene (VIII). This compound was further trans formed into 9 methylphenanthrene, which was identical with a synthetic product,
(I) is the only known alkaloid which does not

contain a beterocyclic N-atom yet is highly poisonous This may be connected with the fact that Colchicum autumnale belongs to the monocotyledons usually devoid of any alkaloidal

constituents. For tetrahydrocolchicine, see D.R.P. 279999, for octahydrocolchicine, C₁₁H₃₅O₂N, see Windaus (Annalen, 1924, 439, 74). (II) does not occur naturally; if obtained by extraction of col-

chicum, it is a secondary degradation product.

Physiological Action of Colchicum -Colchicum is employed in medicine chiefly as a remedy in gout and rheumatism, and colchicine itself, mostly as the salicylate, has been similarly used The alkaloid is highly poisonous and not more than 3-4 mg of crystalline (1) per diem should be given; lethal dose for rabbits 4 mg., for mice 3 mg, for dogs and cats 1 mg. per kg. of bodyweight. (I) is a alow poison which only acts after an incubation period, very aimilar to the one of bacterial toxins. Owing to its slow nervous system, no symptoms are shown for colloid accence still continues to be very rapid,

coloration with ferric chloride characteristic for several hours after its administration. It first excites and then paralyses the nerve codings (1) causes acute intestinal pain with nausea and diarrhoea, and in mammals poisoned with (I) the alimeotary canal ahowa all the aymptoma of acuto gastro-enteritis.

Cf. Winterstein, "Die Alkaloide," Berlin, 1927, p. 147; and Henry, "Plant Alkaloids," 2nd ed., p. 391.

Schl.

COLCOTHAR (syn. Caput mortuum). The reddish brown residue of ferric oxide formerly obtained in the manufacture of fuming oil of vitriol from iron sulphate.

COLEMANITE. A hydrated calcium borate, Ca, B₄O₁₁,5H₂O, containing 50 9% beron tri-oxide. It is found in California at Death Valley in Inyo Co., and in the Calico district in San Bernardino Co.; and although not discovered untd 1882 and 1883, sa now of some importance as a material for the preparation of borax and boracte acid, Cavities in the massive, white material are lined with beautiful, water-clear crystala with a profusion of brilliant faceta, these are monochinic, with a perfect, pearly cleavage parallel to the plane of symmetry, They are considerably harder (II. 4-4) than borax, apg. 242. The mineral forms, together with other boratea and borosilicates, a bed 7 to 19 ft. in thickness in aandstones and

clays.

The compact minerals, priceite from Oregon and pandermite from Asia Minor, have very nearly the same composition

(Ca,B,,O,,,9H,O),

and they have been regarded as impure, massive varieties of colemanite, E. S. Larsen (1917) has, however, hy an axamination of the optical and pandermite, and proved them to be tricking and distinct from colemanite. These also are mined as borates.

L, J. S COLLARGOL. Colloidal compound of alhumen and silver, containing 70% or more Ag COLLIDINES. Trimethylpyridines are ac-

called, e g. 1:3.5 and 1:3.4.

COLLIN. A preparation made by heating an extract of hide powder or gelatine solution with caustic soda and neutralising with acetic acid. Used in the acalysis of tannin and tanning materials (Parker and Pay e, J.S C.1 1901, 23, 648; Wood and Trotman, ibid. 1071)

COLLIBON. Colloidal iron, 10%, with a trace of copper, used for treatment of anamias (Exans Sons, Lescher, and Webb, Liverpool) B P.C

COLLODION (syn. Collodium) v. CELLULOID PLASTICS

COLLOGION COTTON v. CELLULOSE LACQUERS

COLLOIDS. As by far the greater part of matter, both organic and inorganic, with which man is normally familiar is in the colloid state, there is hardly a branch of science or technology where colloid cuncepts have not been applied, absorption (possibly by the gradual formation and hence the literature of the subject is both of an exidation derivative) into the central voluminous and diverse. The development of and several journals are devoted to it alone, while papers on colloid science appear in all the general journals of pure and applied

chemistry.

Colloidal sols of gold, ferric hydroxide, etc., were known by the middle of the last century, and Faraday with characteristic perspicacity attributed the properties of a gold sol prepared by him to the metal being in an exceedingly fine state of subdivision. In 1861 Graham showed that substances which are normally easy to crystallise are characterised by a high rate of diffusion and by the power to pass through certain animal or vegetable membranes; such substances, which included the common electrolytes as well as sugars, etc., he termed crystalloids. A further class of materials, including starch, soaps, many dyes, gums, proteins, etc., which are difficult to obtain in a crystalline form, were found to he characterised by a low rate of diffusion and inability to pass through membranes; these substances were termed colloids. This separation of colloids from crystalloids by dialysis, a process depending on the inability of the former to pass through parchment or collodion membranes, is still widely used.

The term colloid is now employed in quite a different sense from that in which Graham used it, and it is to he considered as referring not to a special class of chemical compounds (although proteins, many dyes, etc., normally form colloids spontaneously) but to a state in which all compounds and elements are capable of existing. From our present standpoint colloids are treated as special examples of two-component systems in which one substance (the disperse phase) is distributed uniformly in the other (dispersion medium or continuous phase). Such dispersed systems can he classified into three types according to the fineness of division or degree of dispersion of the disperse phase: (a) mechanical suspensions, (b) colloidal solutions, (c) molecular

solutions.

The smallest particles which are visible under the microscope are of the order of about 0.1μ , while the average molecular size is about $0.1 \text{ m}\mu$. Between these limits lies the region of colloidal particles, a substance thus heing strictly defined as being in the colloidal state if its particles of size 1-100 m μ are distributed in a continuous medium. It should be realised that there is no clearly defined line of demarcation between the three groups of disperse systems, the colloids being differentiated for practical purposes by their physical properties form a continuously graded series.

The colloidal state is characterised by the predominating influence of surface. The enormous increase in surface area on subdivision of a cube of unit dimensions is given in the table at the

top of next column (Wo. Ostwald).

As subdivision increases to molecular dimensions, intermolecular and interionic forces become of primary importance, so that the effect of surface and hence of adsorption is greatest in the middle of the colloidal range, in the so-called "zone of maximum colloidality." The term colloid should be extended to include all substances (such as charcoal, silica gel, pyrophoric generalised to include all kinds of dispersion

Length of		
side of	No. of	
cubes in cms.	cubes	Total area.
11	1	6 cm. ²
10-1	103	60 cm. ²
10-2	10 ⁶	600 cm.2) region of coarse
	0	> suspensions,
10^{-3} $10^{-4} = 1 \mu$	109	6,000 cm.2) emulsions, etc.
$10^{-4} = 1 \mu$	10^{12}	6 m.2 upper limit of
		Brownlan
10-5 01	1015	movement.
$ \begin{vmatrix} 10^{-5} = 0.1 & \mu \\ 10^{-6} = 0.01 & \mu \\ 10^{-7} = 0.001 & \mu \end{vmatrix} $	1018	${600 \text{ m.}^{2}\atop 6,000 \text{ m.}^{2}}$ colloidal dimensions.
$10^{\circ} = 0.01 \mu$	10 ²¹	c ooo m 2 Sions.
10 =0.001 μ	10	0,000 m J
$=1 \text{ m}\mu.$ $10^{-8}=0.1 \text{ m}\mu$	1024	60,000 m.2 molecular dimen-
1	10	sions.
=1A		Sions.
	•	()

metals) whose surface properties predominate over those due to chemical constitution.

The complete list of possible two-component disperse systems is given below, the dispersion medium being placed first, followed by the disperse phase:

(I) Solid+solid (ruhy glass, alloys).

(2) Solid+liquid (solid emulsions, minerals with sealed liquids).

(3) Solid+gas (solid foams, porous solids).

(4) Liquid+solid (suspensions). (5) Liquid+liquid (emulsions).

(6) Liquid+gas (foams and froths). (7) Gas+solid (smoke, cosmic dust).

(8) Gas+liquid (mist, clouds).

Although it is possible to obtain all the above in colloidal dimensions, Nos. 4 and 5 havo received by far the greater portion of study and will alone he considered here. Colloidal suspensions whether of solid particles or liquid droplets are termed sols.

CLASSIFICATION OF COLLOIDS.—Colloidal sols differ widely in properties hut fall into two hroad classes, of which colloidal gold and gelatine respectively are prototypes. Tho chief differ-ences which serve to classify the two groups of colloids are as follows:

Gold sol type.

Artificial inorganic colloids.

Viscosity almost the same as that water.

Concentration always

Sensitive to small concentration of electrolytes.

Do not swell, unsolvated. Coagulate to form

coarse granules, irreversible.

Gelatine sol type.

Natural complex suhstances.

High viscosity.

High concentration

possible.
"Salted out" only by high electrolyte concentration.

Highly solvated, considerable swelling. Coagulate to jelly, generally reversible.

It must be realised that the ideal examples of the two types are extreme cases of the diversity of colloid properties and that there exist numerous colloids (e.g. the hydrous oxides and hydroxides) whose properties are intermediate. The most prevalent usage at the moment terms the two classes of colloids hydrophobic and hydrophilic according to whether they "hate" or "love" the dispersion medium-water-or lyophobic and lyophilic when the classification is

250 COLLOIDS.

does not disperse spontaneously and is easily scparated (e.g. hy electeolytes) from the dis-persion medium. Gelatine, on the other hand, is hydrophilic es it disperses apontaneously, is stable and, at the same time, becomes highly hydrated. Another type of classification terms the first class of colloids suspensoid, the dispersed particles being solid, and the second class emulsoid, it being assumed, chiefly by analogy with coarser emplaions and from viscosity considerations, that gelatine and other similar substances exist in the form of hand droplets when dispersed in water. There is much to be and for this theory, but it should be pointed out that mercury and certain ether emulsoid self. 1823, 4, 161; 1829, 6, 161; Edin New Phil. (e.g. dilute oil emulsions) are of the gold type, Journ 1828, 5, 328, 1830, 8, 41] It was soon and that in any case the distinction between solid and liquid tends to disappear in the colloidal range where surface tension effects become allimportant. The chief distinction between the two main classes of colloids is almost certainly due to solvation,

COLLOIDS AND THE ULTRAVICEOSCOPE -The heterogeneity of colloidal sols is clearly shown by the Tyndall effect, for though the sol may appear to be perfectly eleer by transmitted light, when a strong beam of light is concentrated in the sol, its path is marked by an opalescence and the resulting light is polarised. This effect is caused by the scattering of light by the dis-persed particles of the sol. It should be pointed out here that solutions of crystalloids (e.g. sugar) also give a faint Tyndall cone, especially when examined by ultra-violet light, and that even the purest water is not completely optically void. When the Tyndall cone is brought to a focus within the sol and the smell volume, thus brilliantly illuminated, examined through a microscope from above, the light scattered retrically by the sol particles is observed in the microscope. This is the principle of the original slit ultramicroscope of Zaigmondy and Siedentopf (Ann. Physique, 1903, [1v], 10, 1), the optical arrangements of which are shown below

Since no direct light enters the miccoscope, the scattering particles appear as a number of bright spots of light against a dark backof origin spots of igns agains a unit occawise the fine suspension cognitive sysground; it is, of course, not possible to see Inneously, owing to the high cost of running and
the actual colloid particles by the ultramaintenance, colloid mills are not an unqualified
by cach particle. The lower limit of the ultracolloid mills, see F. M. Travis, "Mechanmicroscopo is 5-10 m_{ps}, on that highly dispersed chemistry," New York, 1028, where the practical

media. Gold forms a hydeophebic sol since it | the difference of refractive index between the two phases of the disperse system, and so highly solvated colloids such as gelatine in water are difficult to examine by this method. Despite these limitations the ultramicrescope has proved to be one of the most powerful weapons of the colloid chemist. Many optical modifications of the ultramicroscope have been introduced

The most striking feature of the ultramicroscope image is the rapid and unreasing motion of the particles-a phenomenon similar to the Brownsan movement exhibited by small particles such as pollen grains, gamboge, oil droplets, etc., in water, which is visible in the realised that this movement was due to the impact of melecoles on the colloidal particles, impacts on all sides not cancelling instan tancously, and that a visual representation of molecular motion in liquids as pictured by the kinetic theory was thus obtained, quency of motion increases with degree of dispersion. It has been deduced theoretically by Einstein and Smoluchowski and demonstrated experimentally by Svedberg that colloid particles possess the same kinetic energy es gas or dis solved molecules, and ere subject to the same laws (see osmotic pressure, below) Beownian movement, celculations of the rate of movement of molecular particles have been mede which ere in good egreement with the value as deduced from the kinetic theory , the value of the Avogadro constant N has also been calculated by Perrin, who obtained the value N=65×10²³ for gum mastic particles of radius

65 µ THE PREPARATION OF COLLOIDS -As the colloids I state is intermediate between mechanical suspensions and molecular solutions, colloidel sols may be prepared either by dispersion of the former or condensation of the latter. Of dispersion methods the simplest is the spontaneous dispersion of natural colloids, such as geletine or soap in water, cotton in cuprammonium hydroxide and the formation of organosols of nitro- or acetyl cellulose, rubber, resins, etc. in organic liquids. Mechanical dispersion has re-ceived a good deal of industrial attention especially since the advent of the colloid mill. In one of the commonest types of mill the substance to be dispersed, mixed with the disperson medium, is made to pass between two metal discs, placed close to one another and rotating at very high speeds in opposite directions (or between a stator and a cotor). In this process the coarse suspension particles are tomso as to yield particles appeaaching colloid dimenaions. It is generally necessary to have a dispersong agent present in the initial mixture, otherwase the fine suspension coagulates sponsols of definitely colloidal characteristics appear applications of fine suspensions are discussed, homogeneous when examined by this means. Liquid-liquid systems can be easily dispersed. The intensity of the scattered light depends on incerhaincally, but, except where very dilute

has to he added. Electrodispersion was first introduced hy Bredig (Z. Elektrochem. 1898, 4, 514) who prepared metallic sols hy arcing wires of the metal beneath the surface of the dispersion medium. Svedherg has modified this method hy enclosing the wires in a silica tuhe which has a hole hored in front of the arc. The metal vapour produced in the arc is blown out hy a stream of inert gas and condenses to a sol in the dispersion medium, which is thus not subjected to the decomposing influence of the arc. The high frequency oscillatory are has also heen used with great success to produce highly dispersed sols of practically all the metals and hy a modification of the method ether sols of the alkali metals were prepared. Dispersions of solids in gases can he produced explosively hy condensing the gas under pressure in the pores of a solid and then releasing the pressure (see Sata, Bull. Chem. Soc. Japan, 1936, 11, 443). Volcanic dust is produced in this way. One of the most important dispersion methods is that of peptisation, in which a coarse suspension is dispersed to a sol by the action of an electrolyte; simple examples are the peptisation of arsenic sulphide hy hydrogen sulphide and of ferric hydroxide by a small quantity of ferric chloride.

Condensation methods of dispersion are more numerous. Amongst the simpler methods may he mentioned solubility reduction on cooling (e.g. sulphur in alcohol) and the effect of solvent change (e.g. hy dropping an alcoholic solution of gum mastic into water), while the formation of organosols may he accomplished by the simultaneous condensation of a metal vapour with that of its organic dispersion medium (Roginski and Schalnikov, Kolloid-Z. 1927, 43, 67) the applications of this method are limited by the fact that the metal must he capable of subliming at a temperature at which the liquid does not char. Another interesting method makes use of the silent electric discharge (Miyamoto, J. Chem. Soc. Japan, 1934, 55, 1273) wherehy colloidal arsenic can he prepared from arsine and hydrogen. By far the most important condensation method of colloid formation is that of carefully controlled chemical reaction. It has been established by von Weimarn that precipitation from either very dilute or very concentrated solution yields a colloidal product, whereas at medium con-centrations normal precipitates are obtained; excellent examples are afforded by the precipitation of barium sulphate and of Prussian hlue. The formation of colloidal materials at high concentrations is at first sight unexpected, hut the explanation is presumably that, formed in such close proximity, the molecules have no time to arrange themselves into a regular crystalline lattice, hut remain in an amorphous condition. This is horne out hy the fact that these amorphous, high concentration sols are unstable and age with the formation of normal crystalline aggregates. Nearly every type of reaction in solution can thus he made to produce colloidal sols, amongst the most useful may he mentioned the following:

emulsions aro required, an emulsifying agent and numerous organic reducing agents), oxidation (e.g. oxidation of hydrogen sulphide to give a sulphur sol), hydrolysis (formation of Fe(OH)3 sol by hydrolysis of FeCl₃; silica gel), double decomposition (formation of sols of acids, hases, and salts), polymerisation (vinyl esters, styrene, etc., Staudinger, Ber. 1926, 59 [B] 3019). To show the wide range of possible reactions, mention may he made of the formation of colloidal sodium chloride hy double decomposition of thionyl chloride and sodium salicylate and the production of colloidal palladium hydride hy the adsorption of hydrogen hy a palladium sol (prepared hy reduction); the hydride sol is an excellent catalytic reducing agent. For practical details of colloid preparation, see Svedherg, "Formation of Colloids" (1921); Holmes, "Lahoratory Manual of Colloid Chemistry" (1927); Reitstötter, "Die Herstellung kolloider Lösungen anorganischer Stoffe " (1927).

THE PURIFICATION OF COLLOIDS.—Dialysis, the classical method of separating sols from electrolytes is still widely used, dialysers of parchment or collodion, which are constructed so as to have as large a surface as possible, being the most efficient. As this process, depending entirely on diffusion, is somewhat slow, the method of electrodialysis has heen introduced, in which the electrolyte ions are made to migrate away from the sol on the passage of an electric current. The apparatus for electrodialysis is shown schematically below:

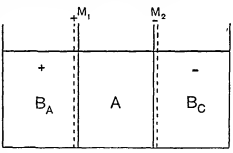


Fig. 2.

The sol is enclosed in the compartment A, separated from BA and Bc hy the dialysis membranes M₁ and M₂, close to which the elcctrodes (dotted lines) are placed. The anode should he of platinum, the cathode may he of copper or any other suitable metal. There are three effects contributing to the separation of ions from the colloid: (a) as in the case of ordinary dialysis, diffusion, (b) migration of ions through the membranes to the electrodes, (c) electro-osmotic streaming of the liquid. In order that the third cause should be effective, the anode membrane should he positively charged (prepared from alhuminoid materials) and the cathode membrane negatively charged (collodion or parchment).

Substances in colloidal suspension pass readily through filter papers, hut, if the close relationship between colloids and mechanical suspensions is horne in mind, it will he realised that reduction (preparation of if the pore size is sufficiently reduced, it ought metallic sols hy hydrogen, carhon monoxide, to he possible to retain colloids hy filtration,

while allowing water as well as dissolved ions concentration reduces the surface tension, U and molecules to pass through. This has been becomes positive; when it increases the auritace successfully carried out by using gelations tension U in negative, indicating a concentration materials, the pures of which are of colloidal in the bulk of the liquid away from the surface dimensions end can be varied in size by model. This negative adorption should obviously be eations in the methods of preparation. The of a much lesser magnitude than normal adsorpgreatest degree of success has been attained by ultrofilters of collodion supported in the pores of filter paper. As the concentration of the impregnating material is increased, the perseability of the filter decreases, so that a series of graded filters can be prepared which may be employed not only to purify colloids, but to sort out the particles according to their sizes (Bechhold, Z. physikal. Chem. 1907, 60, 257). The pore size of ultrafilters may be calculated from the volume of water forced through unit filter area in unit time by a given pressure. It may be stated that ultrafiltration depends not only on the sieve action of the ultrapores but also on the charge on the filter. Collodion filters are negatively charged in presence surface layer, of water and allow the passage of negative. Gibh's fact. particles more readily than of positivo ones. This renders determination of absolute size by ultrafiltration somewhat doubtful. Electroultrafiltration is a device to speed up ultrafiltration in the same way that electro dialysis quickens dialysis. The relative rates of separation of electrolytes and non-electrolytea by the four methods are given in the following table (E. Heymann, Z. physikal, Chem. 1925, 118, 65), a potential gradient of 40 v./cm. being used in the electrical processes :

Process					NaCI	Cane sugar
Dialyas					1	03
Electrodialysis				·	168	2(1)
Ultrafiltration					14	14
Electro-ultrafilt	rati	on	٠.		162	14

(For a review of the literature on ultrafiltration and an account of the separation of single proteins, rubber latex, acetone collodion, etc. by this means, see Ferry, Chem. Rev., June, 1936.)

ADSORPTION .- The enormous surface of contact between disperse phase and dispersion medium possessed by aubstances dispersed colloidally has already been stressed. This is chiefly important because of the ions or molecules which can be attached at the interface by adsorption and thus modify the stability and properties of the system. The term adsorption is here used to indicate a concentration or attachment of some aubstances at the interface. In a two-phase system where one of the phases is e liquid it follows on thermodynamical grounds from surface tension considerations that the most stable relation between the two phases is the one characterised by a minimum surface tension. Hence, if there is present in the hould a substance which lowers its surface tension, this aubstance will accumulate in the aurface layer. The relationship as atrictly

deduced by Gibbs is $U = -\frac{C}{RT} \frac{d\sigma}{dC}$ where U is the

tion, which is in accord with the experimental facts. On the other hand, a relatively minute quantity of a surface or capillary active com-pound will reduce the surface tension enormously. The fatty acids which are particularly effective in reducing the surface tension show an increasing activity with increasing number of carbon atoms in the homologous series (Traube's rule). It is interesting to note also that adsorption of these substances on charcoal increases in the same order. Capillary active substances (eg. the soaps) also act often as emulsifying agents in liquid liquid dispersion ; this property is closely connected with the fact that there is a definite molecular orientation in the adsorbed

Gibh's isotherm cannot be applied directly to solid surfaces where da/dC cannot be measured. The sold surface is always very complicated even with opparently plans crystalline surfaces of metals or saits, there are numerous deep fissures (Smeckel cracks) between the blocks of crystals, as well as peaks and valleys on the aurface. Atoma on peska or edges are less strongly attached to the body of the material and are consequently very active for adsorption and catalysis. Much evidence on active centres has recently been obtained by the study of differential eatalyst poisoning which conclusively demonstrates the existence of different types of surface atoms. Active centres, which are often capable of fixing ions or molecular groups chemically, are of fundamental importance in colloidal behaviour. For solid surfaces, the empiricel sorption isotherm of Freundheh,

 $\frac{x}{m} = kC^{\frac{1}{n}}$ (where x is the amount adsorbed, m the weight of adsorbent, C the equilibrium concentration, and & and n constants), is generally applicable (see, however, J. W. McBam, "The Sorption of Gases and Vapours by Solids,"

London, 1932, p. 5). When an adsorbing surface is placed in contact with an electrolyte solution it can generally be shown that the electrolyte is not adsorbed as a whole, but that one ion is more strongly adsorbed than the other. If, for example, e collord adsorbs the anion preferentially, the excess of cetions will arrange themselves as an outer sonle atmosphere, the complete formation of adsorbed and outer ions constituting an electrical double layer. The colloid surface and nature of the double layer is well illustrated by the precipitation of the alver halides. In the apace lattice of ailver bromide, of which a two dimensional representation is given on Fig. 3a. each anterior salver or bromide son is surrounded by aix equidistant ions of the opposite aign. The ailverions on the surface, however, are only excess of substance at the interface, C its con-centration in the bulk of the liquid, o the surface (three in the plane of the paper, one in front tension, R the gas constant, and T the absolute and one behind), and similarly each surface temperature. It is seen that when increasing bromide inn is surrounded by only five silver

sorbed from solution on to surface silver ions: as the added bromide ions are already halanced hy corresponding potassium ions, a second hut more mobile adsorbed layer (of potassium ions) is formed all over the surface, with the result that an electrical double layer is formed and the particle hecomes a stable negative colloid (see Fig. 3b).

At the point where equivalent amounts of potassium hromide and silver nitrate are present the particles are coagulated as a normal uncharged precipitate, but if an excess of silver nitrate is present a positive sol results, according to the scheme in Fig. 3c. It will be observed that the facts outlined above give an explanation hased on adsorption of the phenomenon of peptisation already mentioned; it is seldom, however, that peptisation adsorption is quite as simple as the case cited here. Further reference to this type of adsorption will he found in the following sections.

ELECTRICAL PROPERTIES OF COLLOIDS .- It has already been mentioned that the particles of a colloidal sol are electrically charged, a property which they share with suspensions and other surfaces. A qualitative determination of the sign of the charge of a colloid may he made hy a simple adsorption experiment, positive colloids

ions; each surface ion thus lacks one ion of the opposite sign to saturate its affinity. When the particle is present in a solution of potassium bromide, bromide ions are preferentially addependent on the physical nature of the compound which forms the sol but, as will be evident from the case of silver hromide already discussed, is influenced by such quantities of electrolytes as may be adsorbed from the

If a direct current is passed through a colloidal sol, the charged particles move towards one or other of the electrodes: this movement is known as cataphoresis and can he measured by the moving houndary method, the houndary heing seen by means of a Tyndall cone; if the colloid is colourless ultra-violet light may be used. The Hittorf transport method can also he applied to cataphoresis, hut hy far the most satisfactory and convenient measurements are made by observing the movement of the actual particles across the field of the ultramicroscope. When the moving particles are discharged at the electrodes, they coagulate, an observation which suggests a close relationship hetween electrical charge and stability, a matter which will be discussed in the next section. The effect of small quantities of electrolyte on the cataphoretic velocity is particularly marked, the influence of the ion of charge opposite to that of the colloid increasing very greatly with valency. If, instead of allowing the particles to move in the electric field, they are held stationary in the form of a diaphragm, the potential difference between disperse phase and dispersion medium manifests itself by a movement of the dispersion medium towards one or other of the electrodes, the direction being the opposite of that in which the diaphragm would move if free. This effect, which is known as electrical endosmose or electro-osmosis, is influenced by the presence of electrolytes, etc., in the same way as cataphoresis. Two other related electrokinetic phenomena have been investigated, namely the streaming potential developed by forcing a liquid through a diaphragm, and the converse sedimentation potential produced by the passage of solid particles through a liquid under gravity. The electrokinetic potential (ζ) is related to the cataphoretic velocity (u) hy the following equation:

$$u = \frac{Dx\zeta}{4\pi\eta}$$

where D is the dielectric constant of the medium, η its viscosity, and x the potential gradient. The velocity of migration of colloidal particles is of the same order as that of ions.

The origin of the electric charge on colloidal particles has been much discussed, one school maintaining that it is due to the adsorption of ions from solution, the other that the charges are produced from the ionogenic substance which forms the disperse phase (see below). It is now almost certain that both causes operate simultaneously in most cases, one or the other predominating with different types of colloid. It is clear also that the charge on the colloid particle is compensated by an equal and opposite (ionic) charge in the liquid in immediate heing adsorbed hy negative adsorbents such as contact with it, the system as a whole being

(like an electrolyte solution) uncharged. The taking place and that colloidal character is whole system of colloid particle plus compen-sating double layer (gegenions) is often termed crete fatty acid anion in solution, but an a micelle. On passing an electric current, the aggregate of fatty acid ions together with two parts of the double layer tend to move in undissociated salt molecules; the colloidal opposite directions, thus setting up an opposing aggregate is really a giant polyvalent ion dise m.f. The intermicellar haud which is a con sociated at the surface and compensated by an ductor tends to compensate this, so that there equivalent number of acdium ions in solution is a net movement of the mucelle towards the im close proximity to the giant ion. When an electrode. With organosols, where the liquid is electric current is passed, the outer sodium ions a non-conductor, the disturbances must be move to the cathode, the giant ions to the anode. electronic rather than ionic, but it should be pointed out that the presence of a very small trace of water plays an important part here. That lassical theory of Helmholtz assumed the

double layer to be compact, there being an abrupt potential gradient between the two phases. This concept has been modified by the diffusa double layer theory mainly due to Gouy (J. Phys. Radium 1910, 9, 457), and it is now conaidered that although the inner layer is probably rigid, the gegenion layer is diffuse and extende to extend the idea of colloidal electrolytes to outwards a distance of a few mit. The propor ordinary sols. In the case of a positive hydroxide tion of positive aggenions (in a negative sol) sol prepared by hydrolysis of ferric chloride, decreases and that of negative ions in the outer the micelle is represented as layer increases with increasing distance from the The presence of electrolyte in the intermicellar liquid influences the diffuse layer, the thickness of which decreases with increasing

concentration of electrolyte.

Some confusion existed as to the relationship between the Nernst electrochemical potential (ε) and the electrokinetic potential (ζ), but Freundlich has succeeded in measuring both potentials for the system glass water, and it is now realised that they do not necessarily bear any relationship to one another. The electrochemical potential represents the total potential drop between the disperse phase and the body of the dispersion medium and is little affected by the presence of ions in the latter. The of the donble layer and the body of the bould. and is thus only a constituent part of the electrochemical potential; it will be smaller Physical Chem 1936, 40, 997) have shown that than the latter, but its insgnitude and sign depend on the ions which are present. The electrokinetic potential thus represents the potential across the plane of shear.

The actual chemical nature of the double layer must now be discussed in more detail. It was placed in water would become negatively layers is improbable but which still display charged owing to a preferential adsorption of electrokinetic properties. the hydroxyl ion ; a basic substance such as ferric oxida would in the asme way adsorb the hydrogen ion preferentially and so become positively charged, the compensating hydroxyl positively cnarged, the compensating nydroxy) (ypo on work as wen as 100 that innory of electrons forming the diffusal aper. With electroly to kneite phenomens, see Abrahamson, Electro solutions a particla will become charged by kneite Phenomens (1933). See also the Farsperferential adsorption of one of the ions in the day Society discussion on celloidal electrolytes manner which has already been illustrated by (Trans. Faraday Soc. 1935, 31, 1) and a useful silver bromide.

the double layer develops directly from a study fuhrung in die Lehre von den Kolloiden." bers it becomes obvious that association is the Brownian movement, and the electric charge

Since the latter contain a large number of undissociated soap molecules in the interior, it follows that some of the sodium migrates to the snode. This and other work by MeBain on soap micelles indicates that, in the case of such definite colloidal electrolytes at least, cataphoresis is a apecial instance of electrolysis Similar considerations apply to acls of most dyes and to many other organic colloids. An attempt has been made, chiefly by Paul and his co workers,

The positive charge is contributed by certain active centres of FeO on the surface, the negative charge of the diffuse layer consisting of an equivalent number of Ci ions Part of the CI is enclosed within the sol particle (as FeOCI) and thus has little influence on the properties of the sol Such colloidal ions may carry several thousand unit charges; the number of chemical molecules (excluding those of the water) per unit charge is termed the colloid equivalent. The classical and modern electrokinetic potential, on the other hand, is colloid equivalent. The classical and modern the potential between the interior, rigid portion theories of electrochemistry are being applied with some success to colloids, regarded as complex electrolytes. McBain and Thomas (J. the transport numbers of ferric oxide sols as well as of the troe colloid electrolytes (soaps, etc.) can be measured by means of the normal (eg Hettorf analytical and moving boundary) methods. But it is difficult to see how the ionogenic theory can be extended to eou alent organic early assumed that a neutral substance when substances in which the formation of ionic double

Electrokmetic measurements on proteins, viruses, bacteria spermatazoa, etc. are yielding valuable results. For a full account of this type of work as well as for tha theory of electrover bromide.

Chapter on the electrical properties of colloidal
The concept of ionogenic substances forming aystems by E. Heymann in Beehhold's "Em-

of soap solutions. The section saits of the lower fatty acids behave as normal lonary idies of a particle in coloni superior are its electrolytes, but in the case of the higher mem.

surface area increases, so does surface energy, and thus, according to thermodynamic principles, colloid particles should have a tendency to aggregation. The rate at which a particle of given size should settle in a liquid under the influence of gravity is given by Stoke's law:

$$V = \frac{2r^2(d_1 - d_2)g}{9\eta}$$

where d_1 and d_2 are the densities of disperse phase and dispersion medium respectively, Vthe velocity of the particle of radius r in liquid of viscosity η . The application of Stoke's law to particles of colloid dimensions shows, for example, that a gold particle of size $10 \text{ m}\mu$ should fall through 1 cm. of water in 29 days, while a benzene sphere of the same size would take about 12 years. Convection and other disturbing influences would normally mask this small rate of settling, so that colloidal solutions in the absence of special influences which encourage coagulation are indefinitely stable. The Brownian movement is manifested as a small diffusion effect superimposed on the external forces of gravity, etc. The Brownian movement, bringing about as it does collisions of the suspended particles, favours aggregation when the impinging particles are uncharged. It is the electric charges on the particles which, by producing mutual repulsion, condition the stability of the sol; uncharged particles of colloidal dimensions, unless protected, coagulate immediately. It has been shown in the case of dilute oil suspensions, that a slow coagulation takes place even when the globules are still feebly charged, indicating that under these conditions only a proportion of the total number of collisions results in coalescence. Coagulation is purely an aggregation effect which is followed by precipitation due to gravity recording to Stelle's leave the initiality appear. according to Stoke's law; the initially amorphous precipitate may then gradually become crystalline.

COAGULATION OF HYDROPHOBIC SOLS.—The effect of electrolytes on the cataphoretic velocity of normal hydrophobic sols has already been mentioned. These systems are influenced by minute amounts of ions, the particles moving more and more slowly as the electrolyte concentration increases until, when the velocity is zero, the sol coagulates, only to be peptised and the particles made to move in the opposite direction if electrolyte addition is continued.

When the colloidal particle is uncharged and does not move in the electric field it is said to be at the isoelectric point, which condition is usually marked by a $p_{\rm H}$ value characteristic of the system. The effect of electrolytes and consequent coagulation of a colloid is an adsorption effect which is due to the antagonistic influence of the separate ions; ions having the same sign as those conditioning the charge of the colloidal particle, tend to increase the electro-kinetic potential, those of opposite sign to decrease it or change the sign. The ions which bring about coagulation are thus those of opposite sign to that of the colloid, and in an

which it carries. On subdivision of a solid, as Although H + and OH - ions are particularly surface area increases, so does surface energy, potent in this connection, the effectiveness of the ion in inducing coagulation increases rapidly with valency, metal ions of the same valency all exerting approximately the same effect. The flocculation values (in millimols. per litre), representing the minimum concentration of various electrolytes necessary to coagulate a negative arsenic sulphide sol, have been shown by Freundlich to be:

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 $\begin{array}{l} {\rm SrCl_2}{=}0.63 \\ {\rm ZnCl_2}{=}0.68 \\ {\rm AlCl_3}{=}0.093 \\ {\rm Ce(NO_3)_3}{=}0.080 \end{array}$ NaCl=51 KCl=50 $\begin{array}{l} \mathrm{KNO_3=}50 \\ \mathrm{MgSO_4}{=}0.81 \end{array}$ $CaCl_2 = 0.65$

The anion has obviously little influence in the coagulation of the negative sol and the flocculation value varies very considerably with the cation valency. When a positive ferrichydroxide sol is investigated, the cation has little effect and the anion valency becomes the predominating factor. Some minimum flocculation concentrations in this case are:

 $\begin{array}{l} {\rm K_2SO_4}{=}0.26 \\ {\rm MgSO_4}{=}0.22 \\ {\rm K_2Cr_2O_7}{=}0.14 \end{array}$ $KNO_2=12$ KCl=9 $\frac{1}{2}$ BaCl₂=9·6

Freundlich (Z. physikal. Chem. 1910, 73, 385) has explained these facts by assuming ions of the same valency to be adsorbed by the colloid to an approximately equal extent. Equal concentrations of any ion of a given valency will thus be required to reduce the electrokinetic potential to zero, and consequently the flocculation value will be the same for all such ions. These concentrations being small, they will appear in the initial part of the adsorption isotherm where the amount adsorbed increases very rapidly with concentration. Supposing that the amount of a univalent ion which has to be adsorbed in order to bring about coagulation is double that of a divalent ion and three times that of a trivalent ion, the steep rise in the adsorption isotherm makes it clear that the actual concentration of salts required to produce this effect is not in the simple ratio 1:1:3, but in the ratio indicated by the experimental results, trivalent ions having a disproportionately large effect. Many ions, especially of the heavy metals and of organic complexes, can be shown by analysis to be more strongly adsorbed than those of the simple ions discussed here, and thus have lower flocculation values than those tabulated above. According to the "chemical" theory of colloidal electrolytes, coagulation is to be regarded as a type of double decomposition; when a ferric hydroxide sol with chlorine as gegenion is coagulated by sodium sulphate, for example, sodium and chlorine ions remain in solution while the sulphate ion is attached to the precipitate.

Although the adsorption theory of coagulation proposed by Freundlich has proved to be a useful working hypothesis, it fails to explain many aspects of the coagulation of hydrophobic colloids. For a complete understanding of this electrolyte solution it is those ions which problem the influence of the coagulating ions on generally have the predominating influence. the sol double layer has to be studied. The

modern point of view is clearly given by Verwey [Congulation in this instance can thus be brought in "Hydrophobic Colloids," a report of a abouthy simultaneous discharge of the stabilising sater symposium held in Utrecht, 1937 (Centen, charge and removal of the stabilising water Amsterdam).

When two colloids sols of the same charge are mixed, there is little notable effect, but when the charges are of different sign coagulation may take place if there is roughly the same amount of each present, otherwise the one which is in excess in part remains dissolved.

The stability of colloidal sols generally decreases with temperature increase, a smaller concentration of electrolyte being necessary to induce coagulation at higher temperatures. Although this effect may often be due to solution of electeolytes from the walls of the containing vessel, results of Reid and Burton (J. Physical Chem, 1928, 32, 425) show that in some cases heat alone may hring about coagulation. Amongst other coagulating infinences may be mentioned mechanical agitation and the effect of radiation of various sorts (ultra-violet light, X-radiations, radioactive rays). The latter effects may often be explained by a photochemical action on protective colloids, peptising electrolytes, etc. Bacterial suspensions may be cosgulated by electrolytes in the same way as ordinary sols.

It is interesting to note that concentrated solutions of electrolytes, and especially of atreng seids, differ so much from dilute solutions that rolloids, far from being coagulated, are maintained in stable suspension in them. Voct (J. Physical Chem. 1936, 40, 307) has prepared sols of morganio compounds in strong adiphune and phosphorie acids and in certain concentrated

selt solutions.

From what has already been said, it is obvious that peptisation is due to adsorption effects of a similar nature to those which cause coagulation of already peptised systems, namely preferential adsorption of positive or of negative ions. Sata (Kolloid Z. 1935, 73, 50) claims to have peptised ferric hydroxide by means of ultrasome waves, even in the absence of electrolytes.

STABILITY OF HYDROPHILIC COLLOIDS .-Hydrophilic colloids are on the whole much more stable than are hydrophobic colloids, although the stability is more dependent on chemical constitution; small amounts of protein decomposition products produced by philie colloids; on the contrary, they have a high hydrolysis on heating may influence the dispersive power, being able to "dissolve" peoperties of the sol considerably. Hydrophile cellulose and other normally insoluble auticolloids at once differentiate themselves from hydeophobic sols by their greater inertness towards electrolytes. The former type is indeed precipitated by a large concentration of electrolyte, but the amount is so much greater that the term "salting out" is used to distinguish the effect from ordinary hydrophohe coagulation. Hydrophile particles are charged in the normal way, the charges being capable of progressive neutralisation by adsorption of Hydrophilic ions to an iso-electric point. colloids do not, however, coagulate at the 150electric point, although this is their point of minimum stability. This increased stability as compared with hydrophobic colloids is due to hydration, the sol particles being covered hy a As the proteins contain both acid and basic water layer like troly dissolved aubstances, groups, they form salts both with bases and

sheath. An important series of observations on this sobject was made by Kruyt (Kolloid Beil. 1926, 28, 1), who showed that a negatively charged agar sol became discharged on the addition of small quantities of electrolyte but was still stable owing to its water sheath. If a dehydrating substance such as alcohol or acetone were now added, the colloid coagulated. On the other band, if a similar amount of dehydrating agent were added hefore the addition of electrolyte, the water sheath was destroyed but the sol remained atable owing to its negative charge. Such a dehydrated hydeophilic sol behaved exactly like a normal hydrophohic colloid, being flocculated by small amounts of electrolyte, the roncentration and valency effects of which were exactly the same as with a negative arsenic

los ebidelus. In view of what has just been said it becomes clear that the "salting out "effect of electrolytes on hydrophilic colloids is due to their dehydrating action as well as to their power of neutralising the charge. As the former effect demands a much larger concentration of electrolyte than the latter, it is apparent that the minimum concentration required to bring about coagulation will depend essentially on the dehydrating power of the salts, there is thus little noticeshie valency effect, in contrast to that of hydrepholic coagulation. When a number of sodium salts are arranged in order of "salting out" efficiency, the following sequence is obtained: citrate > tsrtrste > sulphste > acetate
> chloride > mitrate > chloride > nodide
> thiocyanate. This is the so-called Hofmeister or lyotropic series and is the same as that which represents the power of these electrolytes in "salting out" organic compounds from their solutions and of reducing the solubility of gases in water ; in the Hofmeister series, therefore, the salts are arranged in order of their hydration. In practice sulphates of sodium or magnesium are used for the salting out of colloids because of their high aclubility as well as position in the series. Salts at the end of the Hofmeister acries such as todides and thiocyanates are not capable even in saturated solution of congulating hydroatances.

The proteins are the most important of the naturally occurring hydrophilic colloids; the behaviour of these substances the electrokinetic properties of which are greatly influenced by H+ and OH- and little by other ions, is best interpreted by considering them to be amphoterio electrolytee similar to hut much more complicated than the amino acids.

$$\left(R < \begin{array}{c} NH_a \\ COOH \end{array}\right)_n \rightleftharpoons \left(R < \begin{array}{c} NH_a^+ \\ COO^- \end{array}\right)_n$$

acids. In alkaline solution (e.g. NaOH) the | fore, the smaller the gold number. An alternaacid groups are active and a sodium salt is formed which dissociates to some extent to produce sodium ions and a negative anion. In alkaline solution, therefore, proteins are generally negatively charged and wander to the anode. In the presence of excess acid (e.g. HCI) the hasic groups form chloride derivatives which dissociate to form protein cations and chloride gegenions; the protein is positively charged in these circumstances and wanders to the cathode. There must obviously he some hydrogen ion concentration at which the particles are uncharged or at any rate acidic and hasic to an equal extent; this isoelectric point depends on the relative strength of the acid and basic groups and is thus not necessarily at p_H =7. The isoelectric point is readily determined by cataphoresis, for here the mobility is zero; it can also he measured by precipitation, viscosity, potentiometric titration, etc. There is evidence that at the isoelectric point proteins exist as "zwitterions" (formula b) rather than as neutral molecules (a). The iso-electric point is a specific property of each individual protein and is usually well on the acid side of neutrality, but the isoelectric point of hæmoglobin is at $p_H=7$. Proteins hehave normally in having their minimum stability at the isoelectric point.

PROTECTION.-When SENSITISATION AND minute amounts of a hydrophilic colloid such as gelatine are added to a hydrophobic sol, the stability and cataphoretic velocity of the latter are reduced although actual coagulation may not occur. This effect is probably due to the normal adsorption of the sensitising particles which behave as colloidal electrolytes; tho charge on the hydrophobic sol is thus reduced and unstability results. It is found that sensitisation only occurs when the protein film is incomplete, and the ingenious idea has been put forward that it may thus he due to simultaneous adsorption of the hydrophilic colloid particle by two or more hydrophobic particles which are thus brought into close contact. When sufficient gelatine (or other hydrophilic colloid) is present to form a complete coating on the hydrophobic particles, the latter take on a typical hydrophilic character and are inert to small concentrations of electrolytes; such a system is termed

a protected sol.

A gold sol or an oil emulsion may be flocculated by 0.003% gelatine, although larger, and

much larger amounts stabilise.

The protection of sols is of great importance and has been practised empirically since ancient times. Zsigmondy showed that the sharp colour change from red to hlue displayed hy gold sols under the influence of electrolytes could be used as a means of ohtaining a quantitative comparison of the protective action of different colloids. The colour change supposed to he due to aggregation is inhibited by protection. He defined the *gold number* of a protective colloid as the number of milligrams which are just sufficient to prevent such a colour change on the addition of 1 c.c. of 10% sodium

tive method of measurement is by the rubin number of Wo. Ostwald, in which the weight of colloid in 100 c.c. of solution which will just prevent change of colour in a Congo-rubin sol similar to that of the gold sol is the criterion. The gold and ruhin numbers of a few protective colloids are shown in the table:

Colloid.	Gold number.	Rubin number
Gelatine and glue .	0.005 - 0.01	0.025
Isinglass	0.01 - 0.02	-
Sodium caseinate .	0.01	0.004
Hæmoglohin	0.03 - 0.07	0.008
Alhumin	0.1-0.2	0.020
Sodium oleate	0.4 - 1.0	
Dextrin	6-20	
Potato starch	c. 25	c. 0·2
Silica sol	ø	00

In many hiological systems double or multiple protection is effective; thus in milk, which is essentially an emulsion of fat droplets in water, some workers consider the oil-water interface to he protected hy casein which in turn is protected by the more efficient lactalhumin. The proportion of lactalhumin to casein in cow's, human and ass's milk is 0·175, 1·2, and 3·31 respectively. Of the three varieties, cow's milk is much the most readily coagulated by acids and hy rennin, the function of the latter seems to be to convert lactalbumin by enzyme action to a coagulant. In agreement with the figures quoted the coagulum of cow's milk is much coarser than that of the others. milk is thus much less digestible for an infant than is human or ass's milk but may be protected by the addition of gelatine, dextrin, Irish moss, and other protectors and even by sodium citrate or tartrate, which also possess general protective action.

Protection is essentially an adsorption phenomenon; this is demonstrated by the fact that a finite time interval must elapse after the addition of gelatine to a hydrophobio sol, before protection is complete; further, although the degree of protection is determined by the concentration of protective colloid used, once a hydrophohic sol is protected, dilution does not diminish this effect. It is generally most convenient to assume that the particles of the protected sol are covered by a complete layer of protector or that, at any rate, its active centres are thus shielded, but Zsigmondy has demonstrated ultramicroscopically that, when the particles of hydrophobic sol are small compared with those of the protector, one particle of the latter may adsorh several of the former with the usual protective effects. There is, however, no relationship hetween the sign of the charge of the hydrophohic colloid and protective action, proteins, for example, heing capable of protecting both positive and negative sols. The protected sol assumes the electrical properties of its protector; a quartz suspension protected by gelatine will possess the cata-phoretic velocity and isoelectric point of gelatine.

Protective colloids have considerable industrial importance and there is an extensive chloride solution to 10 c.c. of a standard gold sol. patent literature. As an example may he The greater the efficiency of the protector, there- mentioned the dispersion of colloidal graphite in water (Aquadag) or oil (Orldag) in the interpretation of osmotic pressure results with presence of the protective tannic acid; such preparations have valuable lubricating properties. This process, inveoted by Acheson (J. Franklin Inst. 1907, 164, 375), explains also the ancient Egyptian method of improving bricks by means of infusions of atraw, as mentioned in Exodus, v. (Acheson, J.S C.I. 1910, 29, 246.)

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A further important application is the preparation of reversible metal sols by means of protablic and lysalbic acids, two albumin decomposition products (Pasl and Amberger, Ber. 1904, 37, 124; 1907, 40, 1392). The resulting sols may, in contrast to normal unprotected hydrophobic sols, contain high concentrations of disperse phase, and nn evaporation the protected solid is left as a brittle, lustrous dark-roloured mass which disperses spontaneously in water to reform the metallic sol Protected metal sols such as collargol are now used as mild antiseptics either externally or for intravenous injection.

OSMOTIC PRESSURE OF COLLORDS.-Since there is experimental justification for assuming that the gas laws can be applied to colloidal systems, it should be possible to calculate camotic

pressure from the normal equation $P = RT \frac{n}{N}$ where n is the number of disperse particles per unit volume and N Avogadro's number. As the factor n is inversely proportional to the size of the particles, it is obvious that the osmotic pressure of a sol should increase with degree of dispersion. With hydrophobic sols N is large dispersion. With bydrophobre sols N is large of the membrane, thus replacing the original Na compared with n and the value of the common lond of the congored. This phenomenon is known pressure correspondingly small; thus with a las membrane bydrolysis. If there is a solution of pressure correspondingly small; thus with a as membrane bydrolysis. If there is a solution of typical, dislysed gold sol the cametic pressure sodium chloride on the other side of the memmay be only a few mm, or cm, of water and the corresponding freezing-point depression or rais-ing of boiling-point undetectable. Calculations of the apparent molecular weights of colloid particles by this means are rarely practicable. Osmotic pressure measurements on bydrophobic sols are further invalidated by the relatively large influence of foreign substances and especially of active electrolytes, as the value of a io the equation above represents the sum total of particles (some, molecular, or colloidal) present. According to the micellar throry which we have already discussed, ions are a necessary factor for the stability of colloid systems and so even when all impurities are removed by dialysis the osmotie effect is not due entirely to the colloid particles.
Where allowance has been made for the ionio part of the micelle, reproducible results have been obtained in some cases. The most im-portant work on this subject is that of W. Biltz (Z. physikal. Chem. 1910, 68, 357) and of Duclaux (J. Chim. phys. 1909, 7, 403), who was one of the early champions of the micelle theory. In the dialysis of bydrophobic sols the osmotic effect is often noticeable as an absorp-

hydrophilic colloids are the effect of electroly tes. the presence of decomposition products (of proteins, etc), a ewelling pressure due to solvation and a lack of knowledge as to the degree and uniformity of dispersion. Except in a few special cases there is no reason for supposing that the sol particles are all of the same megnitude and hence the " physical " molecular weight calculated from osmotic pressure data is

only an average value. If, as is generally the case, osmotic pressure is measured by means of a semi-permeable membrane, yet enother complication may be intro-duced by the so-called Doonan equilibrium effects, due to uneven electrolyte distribution. This membrane equilibrium phenomenon was worked out in the first instance for true colloidal electrolytes such as congo red, one of whose ions can penetrate the membrane while the other rannot, but, as the micellar hypothesis suggests, the treatment can be extended to colloidal sols in general. If congo red sol be regarded as consisting of giant rolloidal iona which are incapable of diffusing through negative congo red membrane and of equal numbers of mobile Na + sons, then if there is pure water on the other side of the membrane, each Na + ion which penetrates will cause a molecule of water to dissociate so as to provide a compensating QH ion, the corresponding H + 10n tending to diffuse through hrane from the congo red sol, some of the salt will diffuse through to the sol, but the concentration of sodium chloride will be less there than outside. The actual concentrations on either side of the membrane can be calculated by thermodynamical reasoning, it being assumed that the activities of the diffusible components are the same on both sides of the membrane. This unequal electrolyte distribution gives rise to a potential difference—the membrane potential—and there is an esmotic pressura difference between the two compartments. A considerable number of measurements of membrane potentials have been made which agree fairly well with the calculated results. For a full discussion of this subject, see Bolam, "The Donnan Equilibria," 1932.

THE SIZE AND SHAPE OF COLLOIDAL PAR-TICLES,-It is not possible to measure the size nf a colloidal particle directly in the ultramicroscope, but an approximate measure may be made by rounting the number of particles present at a given time in the field. The volume of this amall portion of sol can be calculated and, knowing the concentration of the tion of water within the dialyser although here sol, it is a aimple matter to determine the average again the effect may be due to electrolytes. In size of the particles assuming that they are hydrophilic sols, the particles are usually of spherical nr possess some other simple shape smaller size than those of bydrophobic sols, and that the density of the dispersed substance hece n is larger and the camotic pressure is the ame as in the solul state. When the sol geoerally measured be although still too small particles are too small to be seen in the ultra-to make freezing-point depressions significant. microscope, Zagmondy's nuclear method (Z. The chief difficulties and inaccuracies in the physical Chem. 1906, 58, 65) may sometimes

be applied. In this method advantage is taken of the fact that in the preparation of, for example, a gold sol, formation of the dispersed particles by reduction requires a finite time during which the solution is supersaturated with regard to gold. If a number of amicroscopic gold particles are added, the gold is deposited only on these nuclei, which thereby grow to a size sufficiently large to be counted in the ultramicroscope. Knowing the concentration of gold in the amicroscopic sol, it is thus possible to calculate the size of the original particles. A rough indication of the size of colloidal particles is also obtained by the use of a series of ultrafilters of different pore dimensions as already described. Many other methods have been attempted though generally with little success, but Wo. Ostwald and Quast (Kolloid-Z. 1929, 48, 83, I56) have succeeded in calculating the size of particles of dye suspensions by a diffusion method. They were able to demonstrate the variation of particle size with the solvents used.

The normally prepared colloidal sol contains dispersed particles of widely varying size and is thus termed polydisperse, in contrast to isodisperse sols possessing particles of uniform size. Although the former is by far the most important type in practice, it is often important to prepare isodispersions for theoretical investigation. Here differential ultrafiltration is of considerable use and Zsigmondy's nuclear method can also be used in specific cases. A further method due to Odén, makes use of the fact that the greater the size of the particles the smaller is their stability and the more sensitive arc they to coagulating influences. On addition of a quantity of electrolyte insufficient to coagulate the whole sol, therefore, only the larger particles are removed. By preferential coagulation Oden was able to prepare a series of sulphur sols of different particle size and properties. Perrin succeeded separating a gum mastic sol into a number of isodispersions by centrifuging at different speeds, and in the hands of Oden and T. Svedberg, sedimentation has become by far the most important method of determining particle size and weight of colloid aggregate. This is discussed in the next section.

The shape of colloidal particles has received a good deal of study by Freundlich and his school, who find that they can generally be assigned to one of three classes, namely approxispherical (all of axes length), disc-shaped (platelets) (one axis much shorter than the other two), and rod-shaped (2 axes much shorter than the third). The last two types of particles are double refracting, possessing an anisotropy due to shape rather than an intrinsic molecular anisotropy. Such a particle only produces diffraction in the ultra-microscope if its length lies in the plane of vision at right angles to the incident light; otherwise it is invisible. As the sol particles are in motion the result is visible as a peculiar twinkling effect. Normally the particles are distributed irregularly throughout the liquid, but on stirring they arrange themselves with as is normally the case, or regularly orientated, their long axes parallel to the stream lines so that as with liquid crystals. streaks appear. This type of behaviour is It has been shown that many substances

shown by many dye suspensions, by vanadium pentoxide sol, proteins, etc. Rod and disc particles can be distinguished by means of polarised light.

Several interesting observations have been made as regards the shape of sol particles, one of which is the change of shape which often takes place on keeping; thus the particles of a freshly precipitated ferric oxide sol are spherical, but age to discs with a corresponding change in optical and other properties; the micelles of many other sols change from spherical to the rod shape on standing. Another property which has been explained on this basis is that of red gold sols to turn to blue during coagulation. This was at one time ascribed to aggregation, an explanation which became untenable when it was discovered that gold particles of widely different size all turned blue, although the final blue particles of one sol could be much smaller than the initial red ones of another. It has now been found that the particles of red gold sol are approximately spherical while those of the blue sol are disc-like.

There are several other properties closely connected with the shape and orientation of sol particles, amongst which may be mentioned the phenomenon of stream double refraction. Freshly prepared vanadium pentoxide sol containing spherical particles does not exhibit the property, but after a few days the liquid which gives no image when at rest between crossed Nicols, shows a bright field when stirred; this is known as stream double refraction. The rodlike particles of a vanadium pentoxide sol may also be orientated by an electric current or a magnetic field, the rods arranging themselves in the direction of the current or of the lines of

magnetic force. X-ray diffraction methods have been applied to colloidal systems with some success. It was early found that a reversible protected gold sol when evaporated to dryness yielded a powder which was essentially crystalline, exhibiting the normal gold pattern, and later many actual sols were shown to consist of crystalline particles. It has already been stated that sols prepared by the interaction of concentrated solutions are initially amorphous, but age to crystalline, precipitates. This is a general effect, it having been shown that amorphous particles tend to result whenever formation has been very rapid or temperature, and hence molecular mobility, low. In general the disorderly arrangement of molecules which characterises truly amorphous material (in contrast to the many pseudo-amorphous or microcrystalline materials, such as charcoal) tends to disappear and is replaced by a regular crystalline lattice. Thus, most freshly precipitated metal hydroxides are truly amorphous and exhibit no X-ray diffraction pattern, but on standing for a few hours at room temperature or for a few minutes in boiling water, a characteristic pattern is developed. In fact just as solids can be crystalline or truly amorphous, so can the molecules of liquids be distributed at random,

normally considered amorphous such as rubber that of helix hamocyanin being 5,000,000. The give characteristic X-ray diagrams when stretched. There is a critical temperature above which this change will not take place and which that of sodium cosinate, for example, which just rises with the degree of stretching. See Weiser, "The Hydrous Oxides," 1936; Faraday Society discussion on liquid crystals (Trans. Faraday

Soc. 1933, 29, 881) SEDIMENTATION AND THE ULTRACENTRIFUGE. -According to Stokes' law, all particles of a given size in a well mixed suspension should fall at the same rate under the influence of gravity. Thus each size-group is deposited at a uniform rate and each decrease in the total rate of sedimentation indicates that some size group has just completed its deposition. If the velocity of sedimentation in a liquid of known viscosity is measured, a determination of the size of the suspended particles may be made, and information obtained as to the heterogeneity of the system. Odén (Proc. Roy. Soc. Edin 1916, 32, 219) used an automatic device to measure the secumulation of a sedimenting suspension on the pan of an accurate balance immersed in the system (see also Syedberg, J Amer, Chem Soc 1923, 45, 943).

As sedimentation proceeds, the accumulation of particles on the bottom produces a local increase in osmotic pressure, resulting in a tendency to upward diffusion of the particles so that an equilibrium stata is eventually set up with a gradually decreasing concentration upwards in the absence of disturbing factors such as convection currents, as was shown experimentally by Perrin for gamboga and mastic suspensions. This is similar to the moleeular distribution of the earth's atmosphere, which becomes less densa at high altitudes

These gravity observations are only applies ble to coarse suspensions, but the method has been extended to highly dispersed colloidal sols by the development of Svedberg's ultracentratuge (see numerous papers by Svedberg in J. Amer. Chem. Soc, also Kolloid Z. 1930, 51, 109. Refined centrifugal methods with high speeds of rotation enable tha force applied to tha particles to be varied so that at the maximum of 44,000 r.p.m., it is 110,000 times the force of gravity. Owing to the persistence of vision, an apparently continuous view of the sedimentation can be obtained and the process followed visually or photographically. Only a few ec. of haud are required for a determination, and equilibrium is established rapidly.

The apparent molecular weight of a Faraday cold sol has been shown to be about 1,300,000. Much smaller particles may be measured by means of the ultrarentrifuge, and the most interesting results are those obtained for the proteins, a surprisingly large number of which, including egg and serum albumin, serum glohulm, legumin, hæmocysnins, etc., are isodisperse. Polydisperse proteins include gelstin, generally lowered by the dispersion in it of casein, lactalhumin, fihrinogen, avoglobulm, hydrophilic colloids. etc., all of which are unstable and show varying dispersity with time. Egg albumin has a moleunperson on the time. Legs around has a more cultar weight of 34,500 and other proteins have immuscible in the protein contains molecular weights that are multiples of this a different concentration of disperse phase—a mount (Svedberg, Nature, 1929, 1225, 1271). The phenomenon discovered by the Jong and Kruyt. hemocyanins have very high molecular weights, (Kolloid Z. 1930, 50, 39) and termed by them

ultracentrifugal method has been extended to sols which are on the borderline of true solutions. begins to exhibit colloidal properties, having a molecular weight of 1,380, and even the heavy ions of exerum can be made to form a sediment. The ultracentrifuge will certainly find increasing use in the future in spite of the high cost of installation. Its hiological applications are important, it being found that viruses, etc., possess remark. s bly well defined ultramicroscopic size and shape.

Some Properties of Hydrophilio Sols A great deal of information has been obtained from the study of the viscosity-concentration relationship in disperse systems Einstein deduced the following expression :

$$\eta_s = \eta_0 (1 + 2.5\phi)$$

where η_s is the viscosity of the system, η_0 of the continuous phase, and \$\dispress the proportion by volume of the disperse phase. The viscosity should thus vary linearly with the concentration of the sol and be independent of particla size, For mechanical auspensions, Einstein's law is obeyed fairly well, but for hydrophobic sola tha rise in viscosity is greater than that which the equation predicts, probably indicating that the factor of has, owing to hydration, adsorption, ete, a higher valua than the volume of the actual substance dispersed. The viscosity of hydrophonosols is much greater than that of the hyuro. phohic sols If Einstein's law is to hold, tha volume of the disperse phase in such a system must be enormously greater than that of the dry colloid. This is one of the strongest arguments in favour of the solvation theory of lyophilic colloids which has already been discussed, When the dry lyophilic colloid is placed in the dispersion medium (e.g., gelatina in water) it commences to swell to several times its original volume by taking up water. Such a swollen gel then disperses directly in water if tha temperatura sa raised, the viscosity of the sol generally increasing for some time afterwards, indicating that solvation is still taking place, To account for these facts it is assumed that the total volume of disperse phase has increased many, and in some cases several hundred, times ; many substances (e.g. agar) in concentrations of less than 1% can take up the whole of the dispersion medium to form pseudo-solid gels. This hydration is not entirely due to direct attachment by the colloid molecules; a large part of it consists in the entrainment and immobilisation of molecules of dispersion medium by intertwining crystals and filaments of the gel substance. On the addition of electrolytes to hydrophilic sols, the first small amounts which slso nentrolise the charge decrease the viscosity considerably; further quantities have httle effect. The surface tension of water is

Hydrophdic colloids may, under certain circumstances, be induced to separate into two

the addition of alcohol or other organic or inorganic liquids to protein sols at 50°. As in the case of flocculation, a decrease in charge and in hydration is necessary to effect coacervation; the phenomenon is not readily treated from phase considerations. Coacervation has siderable biological significance. See Bunenherg de Jong, "La Coacervation, les Coacervates et leur Importance en Biologie," Actualités scientifiques et industrielles, Paris, 1936.

Sol-Gel Transformation.—In the broadest sense, a gel is a substance formed by coagulation of a sol, but the term is often restricted to the transparent or opalescent continuous jellies which are formed by the cooling or evaporation of substances such as agar or gelatine, discontinuous gelatinous precipitates such as those of the metallic hydroxides heing termed coagels. It is probable that the distinction between the two types is not fundamental and that in the latter case gelation is initially uniform, for by choosing the correct conditions, uniform jellies of ferric hydroxide, manganese dioxide, etc., can be prepared which, like silica gel, are produced by

coagulation of the sol.

The hydrophilic colloids behave in very varying fashion as regards gelation. Thus gelatine and agar, which are chemically very different (the former being a protein, the latter a complex carbohydrate), both set to gels on cooling; albumin sols do not do this, yet they are irreversibly coagulated on heating to 60° (cooking of eggs). Still another type of behaviour is shown by a dispersion of cellulose nitrate in benzene and amyl acctate, which sets to a reversible gel when the temperature is raised to 70°. In the gelation of hydrophilic colloids of the gelatine type, the temperatures of setting and melting may be widely separated; this hysteresis in the case of agar is as much as 45°. Melting and setting temperatures tend to approach one another when increasing time is allowed for gelation. Gelation is usually preceded by a great increase in the viscosity of the sol and, as there is no break in the viscosity curve, it is generally assumed that the process, unlike crystallisation, is continuous. It may be, however, that the normal increase in viscosity with temperature decrease, combined with the elastic effects which come into play as the gel is formed, may mask any discontinuity. experimental indication of discontinuity has been noticed by Hatschek (Kolloid-Z. 1929, 49, 244) who found that gas bubbles in a setting sol, which are originally spherical, suddenly become lenticular.

There has been an attempt to explain the properties of gels on the assumption of their heing homogeneous, one-phase systems; this hypothesis has the support of Loeb, Pauli, and Porter, but is not wholly satisfactory. In a recent paper, Hatschek (Trans. Faraday Soc. 1936, 32, 787) has shown that the colour of gclatine gels containing cobalt chloride changes

coacervation. This happens, for example, on mechanically retained. Further evidence against the one-phase gel theory comes from the ultra-microscopic resolution of many such systems.

The results of modern research tend to the conclusion that the gel substance consists of submicroscopic fibrils or threads which may be either solid or viscous liquid, amorphous or crystalline in structure, and which are formed by partial coalescence of the highly solvated hydrophilic particles which thereby enmesh relatively large amounts of the dispersion medium. According to this viewpoint there is no phase reversal on gelation, the dispersion medium remaining the continuous phase and the fact that electrolytes diffuse in dilute gels as readily as in pure water becomes easily understandable.

Some interesting work by de Jong (Z. physikal. Chem. 1927, 130, 205) supports the flocculation theory of gelation but shows that the electric charge and hydration of an agar gel are practically the same as that of the sol. The hypothesis has been developed that charge and hydration are not uniformly distributed on the surface of the dispersed material, but are collected in strongly protected or hydrophilic regions, leaving unprotected or hydrophobic centres at which coagulation takes place. Protected portions of the surface are thus left to form the walls of the capillaries of the resulting porous aggregates. It is also postulated that on lowering the temperatures of a sol, a surface re-arrangement takes place, which results in an increased number of hydrophobic centres.

Gelation is a partial coagulation; the process normally continues in the gel, the spongy walls adhering to one another at an increasingly large number of spots, the gel thereby shrinking and squeezing out the water enmeshed in the capillaries. This spontaneous shrinkage of a gel accompanied by exudation of liquid was noted by Graham and termed by him syneresis. Some gels such as silica gel show increased syneresis with rising concentration, while others (gelatine, agar, dyes, etc.) behave in the opposite manner.

It has already been stated that when a hydrophilic substance is placed in water, it swells owing to hydration. The phenomenon of swelling has been erroneously described as the reverse of syneresis, hut the former is due to the taking up of bound water, the latter to the liberation of mechanically held water. While in syneresis the volume of gel+water remains constant, swelling is accompanied by a decrease in total volume, probably due to orientation of the adsorbed water molecules resulting in a closer packing than the random distribution in the normal liquid state. During the swelling of a hydrophilic colloid considerable pressure is produced; this is probably an important factor in the growth of plants.

The effect of electrolytes on the swelling of hydrogels is very marked, different salts behaving according to the position of the anion in from pink to hlue on drying, when the water the lyotropic series; thus citrates, tartrates, content of the gel is still 30%. Gels containing this amount of water or less form no ice on freezing, so it may be concluded that this it to such an extent that the gel disperses readily percentage of water is chemically held, the rest at room temperature. Swelling is usually at a

and hydrogen ions favouring the effect.

Gels are often classified as clastic or rigid, gelatine being a representative of the former and alica gel of the latter type. The distinction is really one of degree, for although gelatine is distinctly more elastic than silica, the latter

vibrates to give a musical note when the con-

taining vessel is struck. and his collaborators, and termed by him thizotropy, is the "liquefaction" of gels by vigorous ahaking, agitation, ultrasome waves, etc. On laying the sol saide, gelation occurs once more, when the thixotropic effect can be tropic solidification of gels can be increased by a mechanical congulation of the particles, this is termed rheopery. The solidification of

rheopectic aubstances is accelerated by ultra-

ferric hydroxide gel and is shown also by pro- the rule holds, the following may be mentioned : teins (eg. gelatine, although the effect is difficult to demonstrate in this system owing to rapid re-gelation), clays, paints, emulsions and protoplasm. Even silica gel may be thixotropic under certain conditions. These effects seem to be due to the type of packing of the sol or gel particles, the shape of which plays an important part. Thixotropic sludges are much used as drilling fluids in petroleum boring, and Freundlich has demonstrated that the properties of quickeands are due to a similar effect, all the quickeands examined containing finite quantities of thixotropic clays.

The action of ultrasonic waves on disperso systems has recently received a good deal of study. Weak ultrasonics seem first to orientate the particles of suspensions or sols if these are rod- or plate-shaped. Secondly, cosgulation of the coarser particles of any shape takes place by accumulation at nodes or antinodes Higher energy ultrasonics may have dispersing effects and are capable of emulsifying liquids, liquefying

thixotropic gels, etc.; this is ascribed to the

collapse of cavitation in the liquids. THE DISPERSE PHASE RULE.—A great deal of work has been devoted recently to the study of equilibria between gels and other colloidal solids and the dispersion medium. Although the solubility of a crystalloid in water is independent of the total amount present, this is not true of the dispersion of colloidal substances. In general the amount of solid substance dispersed or peptised varies with the amount of solid phase present, reaching a maximum for medium quantities. This relationship was termed by Wo. Ostwald the "Bodenkor-

"solubility" of which increases to a maximum particles have been formed but not allowed to with increasing amount of dye to a constant grow, is not yet clear. It is certain, however,

minimum at the isoelectric point, both hydroxyl amount of water and then falls again as the total amount of solid still further increases.

The full aignificance of the disperse phase rule is not yet realised. Several explanations in terms of adsorption have been proposed, but none are sufficiently broad to cover all the phenomena of the type. Investigation of the effect of the total amount of gelatine present on its specific swelling and apontaneous dis-A peculiar gel-sol transformation which has solution in pure water shows that variation is been much studied of late years by Freundlich doe to soluble impurities in the gel which dissolve in water to form a solution, the concentration of which depends on the total amount of solid present; such solutions, as above stated, have a great influence on the degree of swelling and may explain the disperse-phase effect repeated by shaking, etc. The rate of thixe- in this case. Although with quite pure substances it may not always hold, the dispersesuitable gentle movement which indoces a phase rule is of great practical significance with the ordinary materials of industry.

Descriptions of researches on disperse-phase equilibrium will be found in the Kolloid Zeitsonics of the type which coagulate anspensions. achirit from 1927 onwards; amongst the most Thixotropy was first demonstrated with a important investigations of systems in which

> (a) The "dispersion" or peptisation of gels, spontaneously dispersing colloids, roarse suspensions (e.g. BaSO, red lead, charcoal, CaCO, in ammonia, etc.)

> (b) The stabilisation and cataphoretic velocity of suspensions (eg. the migration velocity of charcoal particles increases with increasing electrolyte concentration for a given amount of charcoal, and with increasing amount of charcoal for a constant electrolyte concentration).

(c) Emulsification of oils (the amount of oil emulsified by a given amount of alkali in a given volume of aqueous phase is maximum for medium amounts of oil).

(d) Swelling of gels.

THE EFFECT OF HYDROPHILIC COLLORDS ON SOLUBILITY, CRYSTALLISATION, AND CREMICAL REACTION .- It has been stated above that a protective colloid such as gelation prevents the coagulation of hydrophobic colloids—preserves a high degree of dispersion. Gelatine might be expected to have a similar retarding effect on the aggregation of ions or molecules to form solid precipitates or crystals. In fact autoprotection is the inhibiting effect of hydrophilic sols on their own crystallisation, with the result that a more or less colloidal solid mass is formed; good examples of this are given by the acap compounds and many biochemicals. It is sometimrs possible, under special ronditions, to obtain these aubstances in a crystalline form.

Gelatine and other hydrophilic sols may provent the aggregation of molecules or dispersed particles of normally insoluble substances, thus perrogel," generally translated as solid-phase prodocing an abnormally great superstatutation. The first is applicable also to disperse. This effect has been attitled by Hedges and Henley, J.C.S. 1928, 2139 with lead joide, solid, disperse phase rule is a more suitable name, and many examples have been noticed by the Excellent obedience of this rule is shown by present author. Whether the supersaturation is many dyes (cf. acid dyes and the solid phase due to the prevention of aggregation of molerule, Wo. Ostwald, Kolloid Z. 1936, 76, 291), the cules, or whether a large number of colloidal sols and gels is often ahnormal.

The work of Tamman and his pupils has shown that crystallisation is essentially controlled hy two factors, first the rate of aggregation and orientation of a few ions or molecules to form crystal nuclei, and secondly the rate of growth of these nuclei at the expense of the material still in solution. If the former process is quicker than the latter, the result is a mass of small crystals; if rate of crystal growth is greater than that of nuclei formation, a few large crystals are produced.

The presence of a hydrophilic sol or gel may modify either or hoth of these factors. First, the protective influence of the colloid may prevent growth, and result in the deposition of a mass of microscopic or colloidal crystals, or even in supersaturation. On the other hand, in a gel the crystallising solution is free from mechanical and convection disturbances and so each growing nucleus comes into contact with fresh solution slowly and regularly hy means of diffusion only. In the absence of the protection effect it would be expected to find large, well-formed crystals. This is the case in silica gel which has no protecting properties, Hatschek has produced large crystals of gold by reduction of gold chloride in this medium, and Holmes has grown crystals of copper, lead iodide, etc., of considerable size. This type of crystals growth in gels has probably played a considerable part in the formation of large mineral crystals in silica rocks. Crystallisation in gels usually has the further effect of modifying the crystal habit of the substance. Very often trcclike growths, characteristic of the crystallisation of thin films, are produced. Such, for example, are the numerous varieties of lead trees and spiral crystals described by King and Stuart (J.C.S. 1938, 642). There may also he a tendency towards spherulite formation. This rounding of crystal contours is made use of in the addition of small quantities of gelatine to ice cream with the result that no sharp crystals of ice are formed and the product is smooth to the palate. The protective action of glue on crystal growth is illustrated in practice by the action of this substance as a retarder in the setting of plaster.

A great many ohservations have heen made on the precipitation of insoluble substances hy reaction in gels. It is usual to carry out such experiments hy mixing in a test tube a dilute solution of one of the reactants with the gel before solidification, and afterwards placing a more concentrated solution of the second reactant above the gel. The substance in more concentrated solution diffuses into the gel where double decomposition takes place. It might be assumed that such a method of carrying out a precipitation in the absence of mechanical and convection disturbances, where the reacting substances mix only by diffusion and the insoluble product remains where it is premodifies the course of the precipitation. It is ibid. 1933, 29, 1. found, however, that precipitation reactions in

that the solubility of substances in hydrophilic | capillary tubes and in porous solids are similar to those in gels, and as chemical reaction in gels is the rule in hiological systems the matter

is of considerable importance.

The precipitate formed hy reaction in a gel may consist of an apparently continuous mass, a limited number of large and discrete particles, a cellular structure, or a series of bands or discs of precipitate separated by clear spaces. Precipitates of the last type, known as Liesegang rings, have heen widely studied and are difficult to explain hy means of any one theory. For an authoritative account and bibliography of this phenomenon, see Hedges, "Liesegang Rings

and other Periodic Structures" (1932).

INDUSTRIAL APPLICATIONS.—With the rapid growth and increasing understanding of colloid chemistry, applications of its principles are to he found in practically every hranch of science and industry. As all plant and animal organisms are colloidal in character, hiological, medical, and nutritional sciences have to take account of the discoveries of the colloid chemist. The colloidal properties of humus, clays, zeolites, etc., enable them, hy hase exchange, to hold soluble salts which would otherwise be washed out of the soil and unavailable to plants. All fibres are of organic colloid origin, and thus the textile industries are colloid industries; especially so is the manufacture of artificial silk, in which nature's colloids have to be imitated. Predominantly colloidal materials are also used or prepared in the manufacture of soaps, detergents and wetting agents; dycstuffs, films, cellophane and plastics; synthetic resins, rubber, adhesives, leather, tars, hitumen and asphalt; lubricants, horticultural sprays, paper, paints, bread, butter and margarine; glass, ceramics and cements; absorbents, decolorising media, etc.,

Bibliography.—Wo. Ostwald's "Theoretical and Applied Colloid Chemistry" (Wiley) and Applied Colloid Chemistry" (Wiley) is a stimulating, general introduction to the subject. Hatschek, "Colloids" (Churchill) is more detailed, and Hedges, "Colloids" (Arnold, 1931) gives a non-mathematical treatment of the subject on modern lines. Von "Kolloidik" (Steinkopff, Leipzig, Buzagn, Kolloidik (Steinkopn, Leipzig, 1936) gives a brief, systematic treatment, Bechhold, "Einfuhrung in die Lehre von den Kolloiden" (Steinkopff, 1934) is a collection of clearly-written articles by different authors, and Taylor, "Physical Chemistry," Vol. 2 (Macmillan, 1931) contains a useful section with extensive references. More comprehensive monocomprenensive monographs include Jerome Alexander, "Colloid Chemistry, Theoretical and Applied" (1926–1935), Freundlich, "Kapillarchemie" (Akad. Verlag, Leipzig), Zsigmondy, "Kolloidchemic" (Spamer, Leipzig). Reference should also be made to the "Colloid Symposium Monographs," edited by H. B. Weiser where "Lorenzia" edited by H. B. Weiser, whose "Inorganic Colloid Chemistry," Vols. I, II and III, 1938, and to reports of the Faraday Society discussions on insoluble product remains where it is precipitated, would he ideal for quantitative study, but we have, unfortunately, no means of determining in what way, if at all, the gel ibid. 1936, 32, 1042; "Polymerisation," determining in what way, if at all, the gel ibid. 1936, 32, 3; "Colloid Aspects of Textiles,"

COLLOPHANITE. A colloidal hydrated of supply is the United States, a certain amount phosphate and carbonate of calcium forming an coming from France and Russia. There are important constituent of phosphorites and rock. Year areas of pine trees in India, and there are phosphates. It is amorphous and optically great possibilities of development of the roan isotropic, and forms concretionary masses midstry there. As the trees are exhausted resembling opal or gum in appearance, with a without equivalent afforestation in the United vitreons to resinous lustre and a conchoidal States, the gathering grounds spread further fracture. The material may be opaque or translucent to nearly transparent, and the so that the American rosin is the product of a colour ranges through colourless, white, yellow, brown, and reddish. An earthy variety is oleo-resin or turpentine is collected by what known as monite. Sp gr. 269-277 (of monito is known as the "box" system. The trees are 2·1); hardness 4½; refractive index 1·569— I·612. A. Lacroix (1910) gives the formula as

{z[Ca,(PO,);]+yCaCO,+zH,O \z[(CaF),Ca,(PO,),]+yCaCO,+zH,O

which includes also the variety called fluocollophanue. W T Schaller (1911) gives the formula as 9CaO 3P₂O₅ CaO CO₂·H₂O+nH₂O. The mineral is found in three classes of deposits. in guano deposits, having been first recognised in the guano of Sombrero, West Indies, in pockets of phosphorites; and in the phosphatic nodules and coprolites of sedimentary rocks ranging in age from the Eocene to the Silurian. Material of the last of these modes of occurrence frequently contains fluorine in addition to carbonate and phosphate, and is distinguished as fluocollophanita.

COLLORESIN & CELLULOSE PLASTICS COLOCYNTH. Butter apple. Colocynthis, Pharmacopona"; deleted "British "United States Pharmacopous," XI, 1936. (Fr. Coloquinte; Ger. Koloquinte.)

The dried pulp of the fruit, freed from seeds, of Cstrullus Colocunthes Schrader (Fam Cueurhitacem), a prostrate perennial of the gourd family, inhabiting the warm, dry districts of the Old World-India, Persia, Arabia, Syria, and the African and European abores of the Mediterranean. It is cultivated in Spain and Cyprus.

(Cf. Bentl. a Trim. 114)
The "British Pharmacopous" ilemands that not more than 3% of fixed oil he extracted by petroleum spirit, and that the ash be not more than 8% The two chief varieties imported are Turkey and Spanish colocynth, of which the former is the whiter, the richer in pulp and the more esteemed.

Colocynth was well known to Greek, Roman, and Arabian medicine, and in the form of solid extract enters into many of the purgative pills of modern pharmacy. It is a drastic purgative, rarely used alone. Small quantities of the extract have sometimes proved fatal; the powdered drug irritates mucous membranes. has been isolated. For mactive constituents, see Power and Moore (J C S. 1910, 97, 99).

COLOGNE YELLOW P. CHROMIUM.

COLOPHONY. Colophony, or common rosin, usually termed simply rossn, is the residue obtained by the removal of the essential oil from the oleo resin derived from various species of pine, chiefly Pinus australis, P. tæda, P. pinaster, and P. laricio. The principal source

afield, and fresh species of pine are met with, number of species In the United States the "boxed" in autumn and winter, that is, excavations of characteristic shape are made in the trunks about 8 m. above the ground. These " hoxes " hold from 5 to 10 lb. of the exudation. The bark is cut away for about 3 ft, above the box and the wood is acored with grooves leading to the box The exudation commences about the following March and goes on till about the middle of October. The tapping life of the tree varies from 5 to 8 years.

In France, a different method of collection is used, known as the cups and gutter system. The production is confined to the Landes district and the pots attached to the pine trees is a familiar eight to travellers in that locality, The principal tree used is Pinus pinaster. The oleo resin exudes from the tree from March to October from meisions made with an axe. This incision is known as the "earre" and is kept open by removal of a thin slice every week and 19 gradually extended to about 12 ft from tha ground. Small gutters made of galvanued iron are fixed so as to lead the olao resm into a amali cup or pot nailed to tha trae at a suitabla

The tree used in India is the "chir," Pinus longifolia, and collection is similar to the French method. If the trees are lightly tapped in France or India they may have a tapping life of 30 to 40 years, but if they are "tapped to death" (gemmage à mort) their active life may be only 5 to 6 years, when they are felled for timber.

The oil of turpentine is distilled off from tha oleo resin, and the residual solid matter, which is melted and strained into harrels, is the ordinary

rosin of commerce

Colophony comes into the market graded secording to colour. That sold under Mark A is nearly black, and numerous qualities exist until that marked W.W is "water white" rosm of the finest quality. It is a brittle aubstance with a glassy fracture, soluble in ether, chloroform, light petroleum, acetone and alcohol. It is soluble in most volatile and fixed oils It softens at about 75° and is completely melted at 120°-135°. As it is the The purgative action is and to depend on two cheapent commercial resin, it is never substances, one of which is alkaloidal; neither adulterated. It consists almost entirely of a free acid or mixture of acids, partly in the form of an anhydride. The principal acid is termed absetse acid (q v.) A small amount of esters may be present. The analytical characters of rosm are as follows:

Specific gravity at 15°=1 0451-1 085. Acid valua=150-175. Ester value = 7-20. Iodine value (llühl)-118-128. Unsaponifiable matter=4-9%.

The iodine value determined hy the Wijs process is nearly double that found by the Hübl process. The former gives very variable figures if the conditions, such as time and temperature, are varied, whereas the Hübl value is far less dependent on small changes of conditions, and is therefore the better figure for a standard, as it is for resins generally.

A useful test for rosin in other substances, such as shellac, is the Storch-Morawski reaction. About 0·1 g. of the substance is dissolved in 10 c.c. of hot acetic anhydride; the mixture is allowed to cool and a little sulphuric acid of specific gravity ahout 1.53, is allowed to flow down the side of the test-tuhe. In the presence of 1% or less of rosin, a redviolet-purple colour appears at the junction of the liquids, which spreads to the whole of the sulphuric acid layer.

An even more delicate test for colophony is one based on that originally devised by Halphen. A solution of 1 part of phenol in 2 parts of carbon tetracliloride is prepared. A very small amount of the substance to be tested, or an extract of the same with trichlorethylene, is mixed with a small amount of this solution, and the mixture placed on a white tile adjacent to hut not in contact with it is placed a few drops of a freshly prepared solution of 1 c.c. of bromine in 2 c.c. of carbon tetrachloride. The hromine vapour is allowed to drift over the solution-a wave of the hand is sufficientand in the presence of rosin a pale hlue to strong violet colour appears at the edge of the spot, which slowly spreads. So little as 0.1% of rosin in a mixture can thus be detected, and with 0.5% the colour is intense. This reaction is known as the Halphen-Hicks reaction.

Rosin is used as the hasis of cheap varnishes and of resinate driers in the paint and varnish industry, in the manufacture of cheap soaps, and for sizing paper. It is also used for the manufacture of rosin oil, and for most purposes for which resins are used. Its friahility and lack of resistance to wear and weather in varnishes render it a poor substitute for the harder and hetter class resins. It is used as an adulterant of shellac.

The chemistry of colophony is a question of considerable difficulty, and cannot he said to be entirely settled yet. So much is historical, and a great deal of the older published work has been shown to be erroneous, that only the recent work which is well established need be dealt with. References, however, to some of the more interesting of the historical papers will be found below.

The complexity of the "rosin," or as they are, unfortunately, often termed "resin," acids (which name will be used here, as it has been used in so many publications), together with the fact that rosin from different pine trees contains different acids, and also that methods of treatment or so-called isolation have frequently altered the character of the acids, have made it exceedingly difficult to even compare, much less reconcile, a good deal of one observer's work with that of another.

To day it may be said that the "pine resin

acids" are all fairly closely related, in so far as they all appear to be huilt up on a hydrophenanthrene skeleton, and on the basis of the isoprene rule. The principal acid, which is the main constituent of American rosin, is ahietic acid. This acid is found in both the dextroand the lævo-varieties, and the two acids are probably true optical enantiomorphs—having identical structural constitutions. Of the acids known as pimaric acid, l-pimaric acid is closely related to abietic acid, but in what exact manner is not yet clear.

It is unfortunate that the names l-pimaric and d-pimaric acids have been retained, as d-pimaric acid is certainly not the optical isomeride of l-pimaric acid. The elucidation of the structure of abietic acid, $C_{20}H_{30}O_2$ ($C_{19}H_{29}$ ·COOH), is chiefly due to Ruzicka and his associates. The same is true in regard to dextro-pimaric acid (v. Abietic Acid).

Reference may he made to the following: C. E. Soane, Bibliography (J. Oil. Col. Chem. Assoc. 1922, 5, 320), in which everything of importance up to 1922 is quoted; Ruzicka and his colleagues (Helv. Chim. Acta, 1922 5, 581; 1923, 6, 677, 1077-1096; 1925, 8, 632-636; 1924, 7, 875; 1927, 10, 915; 1931, 14, 545; 1932, 15, 1300, 1289, 915; 1933, 16, 169); and R. D. Haworth (J.C.S. 1932, 1784, 2248, 2717).

Bacon and Ruzicka (Chem. and Ind. 1936 55, 546) have now thrown some doubt on their formula for ahietic acid. Recent work on l-pimaric acid has shown that the possibility is not excluded that conjugation does not exist in ahietic acid, and that therefore the position of the double bonds may require reconsideration.

"COLORADO" CRUCIBLE v. Assay-

COLORAN B7. A highly sulphonated product of a fatty oil used as a wetting-out agent.

COLORIMETERS COLOUR and COMPARATORS. The name "colorimeter" is applied somewhat loosely to any instrument in which the observer has to arrange a colour match in a photometric field, irrespective of the kind of information which is to be derived from this colour match. Strictly, the name colorimeter should be confined to instruments designed to effect measurement of colour, that is, to provide numbers, charac-teristic of each colour tested, by means of which colours may be classified and their relations to one another exhibited; it should not be given to instruments in which the readings give no information about the observed colour, a colour match being used merely as a criterion for the measurement of something else, say the concentration of a solution. Instruments of this kind are better described as colour comparators. Unfortunately, the existence of well-established trade names makes it impracticable to adhere consistently to this distinction in referring to proprietary instruments, but the distinction in function should he borne in mind, and the first section of this article will deal with true colorimeters, instruments whose primary function is the quantita- [well chosen it will be found that the great tive study of colour que colour, while the majority of other atimuli can be matched second part will deal with colour comparation; exactly by a suitable maxture of the three whose primary (and usually only) function is standards. The quantities of these standards the measurement of something other than colour.

Colorimeters -The different colour sensations which light may excite in the normal human eve arise from differences in the spectral distribution of energy in the radiation entering the eye, or, more briefly, in the spectral quality of the stimulus. The theory of the relations between the physical stimuli and the colours observed would carry us outside the scope of the present article; suffice it to say that the essential alternative systems from which selection could properties of stimuli which determine whether or not they will produce a visual match are because of its practical convenience, the found to be functions of three independent quantities of the three standards which are variables. Three independent numbers are required to match a standard white (see later) are necessary to specify, quantitatively, the match | regarded as equal. Measurements are converted ing properties of any stimulus, and any two to this basis by multiplying the observed scale stimuli, whether physically similar in apectral readings by suitable numerical factors, usually composition or not, which have all three termed the units light factors. These factors are numbers equal, will produce visual effects easily obtained from the scale readings observed identical both in colour and brightness. This when matching the standard white fact necessarily has its origin in some triple based on this convention are said to be expressed structure of the receiving mechanism in the intrichromatic units. Hereafter, any reference to eye. Little is known of the physiological cause the scale readings of a trichromatic colors of this triplicity, but the fundamental variables meter will not mean the numbers directly may be regarded as the degrees of stimulation of observed on the scales of the instrument but three independent sets of receptors, differing the equivalent numbers on the trichromatic from one another in their relative sensitivities bases obtained on reduction by the white light to the various parts of the spectrum Differ- factors. ences in the relative degrees of stimulation of the three receptor systems are recognised as differences in colour ; differences in the actual degrees of stimulation, without change in the relative degrees, are recognised as differences in intensity or brightness. Any two stimuli, whatever their spectral composition, which produce the same degrees of stimulation, both relative and actual, in the three receptors, appear identical both in colour and brightness; the eye can analyse no further.

fundamental variables characteristic of any stimulus. Fortunately, it is not necessary to do must be equal to the total quantity of the so in order to measure colour. If the visual light in the comparison field, and the result values of all stimuli in terms of the three fundamental variables were known, it is obvious that merely by solving simultaneous equations these variables could be eliminated and any or all of the stimuli expressed in terms of any three of them. This shows that it is possible to express the visual properties of stimuli in terms of variables which are simply quantities of three other stimuli. This being so, it should be possible to effect direct determination of those quantities without troubling about the fundamental variables, and this is in fact done in the type of trichromatic coefficients of b in terms of the instrument now to be described.

principle of a matching field one side of which is Colour quality as thus defined is independent illuminated by the simulus being measured, the of intensity. It depends on the constitution other side being illuminated by a controllable of the light and not on its amount. It is mixture of three standard stimul, which may possible therefore to have materials exhibiting consist of monochromatic radiations obtained the same colour quality but differing in brightspectroscopically, or may be produced by ness. For instance, a bright yellow and a dark passing white light through colour filters. If brown may have the same colour quality, but the the colours of the three standard stimuli are former may reflect ten to twenty times sa much

used in any match are indicated on the scales of the instrument,

These ecale readings do not, however, afford a convenient comparison of the quantities of the three standards. In comparing quantities of hght of different colours it is desirable to express them in units the significance of which is independent of the peculiarities of construction of any particular instrument. Any choice of units is arbitrary, and there are numerous be made In the system most widely used,

In order to measure the colour of a specimen of material under a given illumination, which may or may not be the standard white light on which the scale factors are based, the light reflected from the specimen, if opaque, is uitroduced to the test field of the colorimeter and matched in colour and brightness by a suitable mixture of the instrument standards Denoting the standards by A. B. and C. and the quantities lentical both in colour and brightness; the control analyse no further.

(reduced acale readings) required to match the specimen, S, by u, L, and c,. The quantity of it is not easy to ascernan the values of these the high from 5 present in the field of twee, when evaluated on the same system as the standards, of this colour match is expressed in the equation ;

 $(a_s+b_s+c_s)S=a_sA+b_sB+c_sC$

Dividing both sides by the total quantity the proportional composition of one trichromatic unit of the light from S is

$$S = \alpha A + \beta B + \nu C$$

where $\alpha = a_s/(a_s + b_s + c_s)$, etc. The quantities a_s \$, and y, whose sum is unity, are termed the strument now to be described.

The Trichromatic Colorimeter consists in they define is termed the colour-quality of S.

of the incident light as the latter. Any two completely define the appearance of materials related in this way can be made to specimen under the kind of illumination for look alike by increasing the intensity of illumination of the darker specimen (or diminishing that of the lighter) until the absolute amounts of light reflected from each are equal. However, objects are not normally compared in this way; our ideas of their appearances are based on comparisons under the same intensity of illumination, and brown is therefore regarded as a different colour from yellow. In order to define the colours of materials completely we must measure not only colour quality but also the property which determines their relative brightnesses. For this purpose it is necessary to adopt a standard of comparison, and the one now adopted in standard practice is the brightness of a magnesium oxide surface, prepared by smoking a silvered plate over a magnesium flame. The brightness of this surface when illuminated at an angle of 45° and viewed normally (or vice versa) closely approximates to that of a perfectly reflecting, perfectly diffusing surface, and is given the value unity. The ratio of the brightness of any material to that of the magnesium oxide standard under the same intensity of illumination is termed the brightness factor of the material. In order to determine the brightness factor we utilise the same observations as for the trichromatic coefficients, but require in addition to make a similar measurement with the magnesium oxide screen in place of the specimen. Denoting tho reduced scale readings for this match by a_m , b_m , and c_m , the following equation expresses the quantity equation in trichromatic units:

$$(a_m+b_m+c_m)\mathbf{M}=a_m\mathbf{A}+b_m\mathbf{B}+c_m\mathbf{C}$$

whilst for the specimen under the same illuminatien it was:

$$(a_{\mathcal{S}}+b_{\mathcal{S}}+c_{\mathcal{S}})S=a_{\mathcal{S}}A+b_{\mathcal{S}}B+c_{\mathcal{S}}C$$

The ratio of the quantities of light reflected from the specimen and the magnesium oxide, expressed in tricbromatic units, is therefore $(a_{\delta}+b_{\delta}+c_{\delta})/(a_{m}+b_{m}+c_{m})$. If brightness factors were only of interest as part of a colour measurement, this would be the most convenient way of evaluating them; but brightness factors are frequently measured by photometric methods for purposes unconnected with colorimetry, and it is desirable that they should always be evaluated on the same basis. In calculating brightness factors from colorimeter measurements it is necessary to obtain the result in ordinary photometric units. Quantities of the three standards which are equal on the trichromatic basis would not be equally bright. It is possible, bowever, to determine, by appropriate experiments the relative brightness, or luminosity, of tricbromatic units of the three standards. If these be in the proportion La: LB: Lc; then it is evident that the brightness factor of the specimen expressed as a photometric ratio is:

$$(a_{\mathcal{S}}\mathbf{L}_{\mathcal{A}} + b_{\mathcal{S}}\mathbf{L}_{\mathcal{B}} + c_{\mathcal{S}}\mathbf{L}_{\mathcal{C}})/(a_{m}\mathbf{L}_{\mathcal{A}} + b_{m}\mathbf{L}_{\mathcal{B}} + c_{m}\mathbf{L}_{\mathcal{C}})$$

The colour quality equation and the brightness factor (usually expressed as a percentage) of the optical system associated with the

which the measurements are made.

If the three standard stimuli of a trichromatio colorimeter are chosen at random it will be found that although many colours can be matched by suitable mixtures of the standards there are others which cannot. This applies to every possible set of standards, but by a suitable choice it is possible to secure that the great majority of the colours of commerce can be directly matched. For this purpose the most suitable standards are saturated colours the hues of which match the hues of the spectrum in the red at wave-length about 0.63μ , in the green at wave-length about 0.53μ, and in the blue at wave-length about 0.455μ .

Even then, however, there are some colours which cannot be matched by any combination of the standards. In such cases it is always possible by adding some one of the standards to the colour under test to obtain a modified colour which can be matched by the other two. The result of this process, when the various quantities are all collected to the appropriate sides of the equation, is to give an expression for the colour quality of the specimen in which one of the trichromatic coefficients has a negative sign, for example,

$$S = \alpha A - \beta B + \gamma C$$
,

where $\alpha-\beta+\gamma=1$.

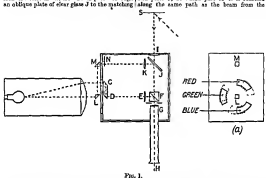
In order to meet such cases and render the instrument universally applicable, it is necessary in constructing a trichromatic colorimeter to make provision for throwing light of any of the three standard colours into the side of the field normally occupied by the test colour alone. Failure to make this provision in the early instruments greatly restricted their utility and prevented them from becoming widely used for scientific or industrial purposes.

The fact that the three instrumental standards are always red, green and blue has given rise to a popular belief that red, green and blue are primary colours, from which all others may be built up. As has been seen, this is an erroneous supposition; there are no three colours from which all others can be produced, though red, green and blue in combination can match a very large proportion of the colours of material objects. A secondary consequence of the general use of red, green and blue standards is that it is customary to denote them in colour quality equations by R, G, and B, instead of by the more general symbols here used.

For many years the only universal trichromatic colorimeter obtainable commercially has been the Guild colorimeter (J. Guild, Trans. Opt. Soc. 1925-26, XXVII, 106). In this instrument the three standards are obtained by passing light from an opal-bulb gas-filled lamp through specially selected red, green and blue filters fitted in annular openings provided with adjustable shutters by which the effective length of each opening can be controlled. light transmitted by these openings is gathered by a rapidly rotating periscopic prism which combines the three coloured beams on the axis

is sufficiently rapid, flicker disappears, and a cube is a aquare aubtending about 2° at the as successful, respectively and as the second of the secon matio view of the essential parts of this colonthe same colour as the red, green or hise
meter. CD is the periscopic prism which rotates standard, which may be required, as already
about the optic axis DE, along which ultimately indicated, when measuring a colour one of whose
passes the light collected by the peripheral end
the promote of the collected of the peripheral end
the passes the light collected by the peripheral end of the periscope as it rotates past the annular is collected by the right-angled prism L, passes windows shown in the end view, Fig. 1a. The to M, which reflects it through a small aperture specimen under test is mounted at S. Light in the wall of the instrument to the lens K from it enters the window I and passes through and plate J, at which part of it is reflected

matching field. When the speed of rotation cube F. The field of view provided by this



specimen S. Between the prism M and the of the surface, remote from the direct unmixed aperture can be fitted one of three filters beams, is directed by a suitable optical system similar to those used for the standards. The to the matching field. This instrument has the light superposed on the test field can thus be advantage that there are no motor-driven given the colour of any one of the standards. parts. It is therefore silent in action and can be An annular photometric wedge N serves to hult on a lighter and more compact scale. The adjust the intensity of the beam to the required provision for determining negative coefficients, amount. This wedge and also the three main and the suxibary equipment generally, are on the shutters are operated by control knobs outside same lines as the corresponding features of the the instrument.

Recently another trichromatic colorimeter. designed by R. Donaldson (Proc. Physical Soc. 19, 98) has described another trichromatic 1935, 47, 1068), has been placed on the market. colorimeter which in principle is identical with The principles of operation are the same as in the Donaldson instrument. the Guild instrument, the main difference in

Guild instrument.

Manfred Richter (Z. Technische Physik, 1938,

A trichromatic colonmeter in which the three the Guld instrument, the main difference in design being in the method of uniting the three design being in the method of uniting the three standard colours. In the Donaldson colormater radiations isolated apectroscopically has been the light entering by the three coloured aperture radiations isolated apectroscopically has been is concentrated by a large condensing less in a Size-28, 29, 292. 393. This type of instrument the inner surface of which is contect with research that the standards are more directly magnetium oxide. The colorum are thoroughly defined in simple physical terms than standards defined in simple physical terms than standards mixed by the multiple reflections to and fre obtained with colour filters and, also, there is inside the sphere, and light from a small region less compensation of certain types of visual

is, however, a disadvantage in applied colorimetry, where it is desirable to eliminate as far as may be practicable the visual peculiarities of the observer. For this class of work instruments with filter standards are preferable. They provide better compensation of minor visual trouble in the hands of semi-skilled observers.

The trichromatic colorimeter, in one form or another, is the fundamental instrument on which all colour measurement is hased. It alone makes direct use of the three-fold relation between the physical stimulus and the observed colour imposed by the characteristics of human vision. The great generality of the method has, however, stood in the way of its adoption for industrial purposes until quite recent years. The fact that a colorimeter could be constructed with any three colours as standards and that no two instruments were ever made alike in this respect meant that results obtained on one instrument were not comparable with those obtained on another. The application of measurement to industrial needs requires that one set of standards shall be universally used. It would be unscientific, and in fact impracticalle, to adopt the standards peculiar to one instrument. Fortunately, there is no need to do this.

The relations between colours exhibited by their values on any one trichromatic system necessarily originate in the relations between the excitations of the visual mechanism by the various stimuli, including the standard stimuli themselves. It follows that the values on systems with different sets of standards must he related, since the same phenomena of perception underlie them all. These relations are of a simple character and make it possible to convert colour values on any one trichromatic system to any other, and we may adopt a set of reference standards which are not necessarily incorporated in any actual instrument and convert the measurements made on any trichromatic colorimeter to this reference system. The results will then be strictly comparable with those obtained on any other trichromatic colorimeter. The theory of such transformations was first outlined by H. E. Ives (J. Franklin Inst. 1915, 180, 673; 1923, 195, 23), and is given more fully by Guild (Trans. Opt. Soc. 1924-25, 26, 95, 139).

The first attempt to introduce this type of standardisation into colorimetry was made in Great Britain in 1925, when the National Physical Laboratory adopted as reference standards the three monochromatic radiations of wave-length 0.700 μ , 0.5461 μ , and 0.4358 μ . To meet the requirements of commerce, systems of measurement must have international recognition. These standards did not meet with universal approval in other countries because of the existence of negative coefficients in the expressions for some colours. As already said, systems with real colour standards. It is,

ahnormality, a point of importance when however, possible to invent imaginary standards the object of the work may he to reveal and which have no real existence, but which are study these ahnormalities. This last feature related to a set of real standards by relations of the same kind as those which connect two sets of real standards. By choosing suitable numerical coefficients in these relations we can define a reference system in which every real colour has all three trichromatic coefficients positive. In 1931 the Colorimetry Committee abnormalities in the majority of practical of the Commission Internationale de l'Éclairage cases and are on the whole less liable to give adopted a standard system of this kind. Because the standards are not real colours, the common notation R, G, B was ahandoned and replaced by X, Y, Z. The recommended symbolic form of the colour quality equation of a stimulus S on this system is

S = xX + yY + zZ

where x+y+z=1. In a practical case, x, y, and z are replaced by the appropriate numerical values. The National Physical Laboratory Report on any trichromatic colorimeter sub-mitted for test gives the constants necessary for converting measurements made with the instrument to the International reference system.

Standard scales of measurement are not in themselves sufficient to ensure unique results for the colour of any given material. The colour of a material cannot be dissociated from the quality of the illumination in which it is viewed. Most coloured objects appear different in daylight and artificial light. The kind of illumination must be specified for every colour measurement. Usually the term colour means colour hy daylight, and comments, such as "Purple objects look red in lamplight," are accepted as reasonable hecause to most people a purple object is one which has that colour by daylight. This is regarded as the real colour of the object and its appearance in another kind of illumination, though it is just as real, is regarded as misleading.

For ordinary purposes of classification colours are measured as seen in daylight, is not, however, a satisfactory light for use in colorimetry; it is neither sufficiently constant in intensity nor in colour. Artificial sources of definite and reproducible spectral composition are required. Further, one such illuminant is not regarded as sufficient to meet the requirements of commerce, and the Commission Internationale has established three, which are designated standard illuminants A, B, and C. The first of these is simply a gas-filled lamp operated at a colour temperature of 2,848°K., and is to be used for the colorimetry of materials intended principally for use indoors in artificial light. The second and third are obtained by combining with this lamp one or other of two blue filters consisting of suitable salts in solution in glass containers. Illuminants B and C are similar in spectral constitution to two typical grades of daylight, one rather yellower and the other rather bluer than the average in this country. For the commercial colorimetry of materials whichever of these three illuminants is most likely to be appropriate should be used, and whichever is used should be specified. this feature is common to all trichromatic Of course, for special purposes, special kinds of illuminant may be called for.

The colours of many materials also depend to a | 1929, 18, 96; 1935, 25, 305) has devised an considerable extent on the angle of medeaco instrument for performing automatically all the of the light and the angle from which they operations involved in the spectrophotometric are viewed. Here again, special cases call analysis of a specimen and the calculations of for special treatment, but it is desirable to have standard conditious which may be assumed to hold unless otherwise stated. The Commission's recommendation is that specimens shall be illuminated at 45 to the surface and viewed method of applying the indirect method to normally to the surface. Where the pecuhar routine colorimetry. Its necessarily high cost is, nature of a material or the nature of a problem under investigation requires departure from these conditions, those actually employed should be stated.

When all conditions of measurement have been specified, there is still uncertainty in colour measurement snsing from the variations of colour vision of observers Despite state ments which have been made to the contrary, any marked abnormality of vision is easily detected, and no person with serious defects of colour vision should be employed in colorimetry There are, however, minor differences in colour vision among observers who connot be classed as defective, and these cause systematic differences in their measurements. One great advantage of the trichromatic method of colorimetry is that when the instruments are used in the proper manner there is considerable compensation of such systematic errors in the majority of practical cases, and the results are much less dependent on the observer than as generally supposed. Nevertheless, differences exist which, if not corrected, would seriously limit the utility of colour standardisation. In addition to the other standards a standard observer is also required, and the Commission has defined such an observer by means of tables showing the results which he would obtain if he measured the various colours of the apectrum. These are based on tables prepared by Guild (Phil Trans. 1931, 230, A, 149) from measurements made by a number of observers in investigations conducted separately hy him and by W. D. Wright (Trans. Gpt. Soc. 1928-29, 30, 141). By means of these tables it is possible to calculate the colorimetric values which would be obtained by the standard observer for any stimulus of known spectral energy distribution. In suitable cases, therefore, the standard values can be obtained for a speci men by calculation. By having a few standard specimens of various colours colibrated in this way an actual observer, by measuring these specimens, can determine the nature and magni tude of the corrections which his observations require.

From the economical point of view it is not feasible to use this method of calculation in all cases and dispenso with direct measurement on colorimeters. The spectrophotometric measurefor a colorimetric measurement. purposes. A. C. Hardy (J. Opt. Soc. Amer. the Commission Internationale de l'Éclairage for

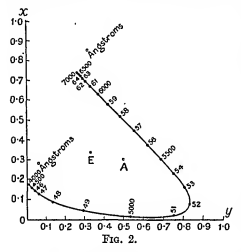
its colorimetric constants for any of the standard illuminants. The machine cannot be described in the space available here. Despite its extreme complexity it appears to afford a satisfactory bowever, an obstacle to its widespread use, and at cannot at present be regarded as an easily available substitute for visual colorimeters.

Another line of development to which attention has been given in recent years is the substitution of the human observer by a system of photo electric receivers. It is possible to design an instrument of this kind which will give directly the values of x, y, and z in the standard colour quality equation of a specimen without the auxiliary calculations required when a visual colorimeter is used. The theory of such an instrument is simple and is given fully in s paper on Colorimetry and its relation to Photometry by Gudd (Trans Illuminating Engineering Soc. London, 1937, 2, 127), but the practical difficulties are great and there is not yet any satisfactory photo-electric colorimeter on the market.

Bleanwhile various instruments, whose indications consist of the responses of a photo cell modified in spectral sensitivity by colour filters, have been put on the market as colonmeters. In every case within the writer's knowledge the claim to this title is unjustified, there being no attempt, successful or otherwise, to incorporato any of the essential properties of a colorimeter. Such apparatus is not, however, without its uses in a colour laboratory. For example, for checking the accuracy of reproduction of a colour for which the same pigment is always used, or for any other test in which identity of colour implies similarity of physical and chemical properties, the responses of a photo-cell used in conjunction with two or more colour filters provide a rapid and accurate means of determining colour equality. But if attempts with such apparatus be made to compare colours prepared from different materials, or even from the same material by different methods of treatment, the results may be most misleading. It cannot be too strongly emphasised that in order to measure colour it is not sufficient to have an instrument the readings of which vary with colour. The readings must depend only on colour and not on other factors. The same readings must always be obtained for colours which are visually identical irrespective of whether the spectral constitutions of the stimuli are similar or not. Further, it must never be possible for the instrument to give the ments required for the indirect method must be same readings for colours which are not visually made with great accuracy, and if earried out on identical. Neither of these conditions can be the usual types of equipment, these measure fulfilled by a physical colorimeter unless the ments, and the subsequent lengthy calculations, spectral sensitivity of the receiver, as modified occupy a time greatly in excess of that required by the various filters employed with it, is I'er routine correctly related to the properties of the eye as work the cost is probibitive and the method can defined by the tables constituting the standard only be profitably employed for standardisation observer. The complete recommendations of

colorimetric standardisation are given in Compte Rendu des Séances, 1931, of the Commission (Cambridge University Press), and also, together with an account of the derivation of the system and instructions for its practical application, in a paper by T. Smith and J. Guild (Trans. Opt. Soc. 1931–32, 33, 73).

From the nature of colour-quality equations on the trichromatic system it is easy to exhibit the relations of colours by a diagram. In the equation $S=xX+yY+\dot{Z}$ we know x+y+z=1, so an explicit statement of the values of any two of these coefficients implies the third and is sufficient to define the colourquality. Colour-qualities may therefore be represented by points on a two-dimensional graph, using some pair of the trichromatic coefficients as co-ordinates. In Fig. 2, for example,



the curved line is the locus of points corresponding to the various spectral colours for the C.I.E. standard observer, using the x and y coefficients of the standard reference system. The point E, for which $x=y=\frac{1}{3}$, represents the colour-quality of an equi-energy spectrum. The point A represents the colour-quality for which x=0.3, y=0.5, and z (by implication)=0.2. A diagram of this kind has many uses. From its geometrical properties we can determine graphically the colour resulting from the additive combination of any group of two or more colours in any proportions. Details of such calculations are given in some of the papers already quoted. The diagram is also useful for studying the changes of colour which take place during the fading of a pigment, or the variation of colour with concentration, and so on. For example, if the colour-qualities of a fugitive pigment at various times after the beginning of exposure to a fading agency be plotted and the points so obtained be joined by a continuous line, the whole course of the fading process can be seen at a glance. It is also useful in such cases to plot a graph of the brightness factor against the other variable—time of fading, concentration, etc.

others the reverse is the case, while with many, both the brightness and colour-quality change. This method of studying colour phenomena has been much utilised in the research laboratories of the colour industries.

Other types of colorimeter employing the principle of additive colour mixture have been described and used for special purposes. In some of these colours are matched by mixing white light with monochromatic light of suitable wave-length, e.g. the colorimeters of P. G. Nutting (Bur. Stand. Bulletin, 1913, No. 9) and I. G. Prest (J. Opt. Soc. Amer. 1924, 8, 173); in others by combining suitable pairs of monochromatic stimuli, as in the Vector colorimeter of Guild (Trans. Opt. Soc. 1925-26, 27, 139), and others which have since been developed on similar principles. All such methods derive their ultimate significance from the fundatrichromatic method, mental and obtained by them can be converted to the standard trichromatic basis, and vice versa. None of these instruments is used in industry, as far as the writer is aware, and details of their construction is omitted from this article.

A somewhat different method of colour measurement, which also in principle depends on the additive mixing of standard stimuli, is employed in the Ostwald system of colorimetry. In this system a colour is expressed by the equation C+W+B=1, where C is the amount of full colour, W the amount of white, and B the amount of black, expressed as proportions, required to match the colour. The "full colours" consist of a series of colours of fairly high saturation ranging in hue from the red of the spectrum through orange, yellow, green and blue to violet, and thence through purple, magenta and cerise back to red. These colours form what Ostwald terms the chromatic circle. A mixture of one of the full colours with white produces various degrees of paleness of colour, while admixture with black reduces the brightness. Important claims are made for this system, not the least of which is that it is the only system of colour specification which takes account of the psychological and æsthetic aspects of colour. It may be stated, however, that these considerations are extraneous to the principles of colour measurement as developed in Great Britain and the United States and embodied in the international system. The subjective relations of dissimilar colours are the proper concern of the user of colour-the artist, architect, etc.—who aims at obtaining certain esthetic effects by his manipulation of colours; but they are not the concern of the colorimetrist, who may only take account of those relations which are necessary and sufficient to provide a basis of classification fulfilling the fundamental requirements of measurement. The only subjective relation which is required, or which is permissible, is the relation involved in colour matching—that of appearing alike both in colour and brightness. For practical measurement on Ostwald's system examples of the standard full colours are necessary. These are generally With some pigments the change in fading is supplied as specimens of dyed wool mounted on almost entirely a change of brightness factor slides convenient for insertion in a suitable with little change of colour-quality; with optical instrument. When the appropriate full

be obtained as accessories to the Zeiss-Pulineh photometer, to he described later in another

In addition to the methods of colorimetry based on the additive combination of standard stimuli already described, colour matching can of something more by each succeeding filter, with corresponding changes of colour. This is termed the subtractive method of colour mixing. Colours formed in this way are in no sense a mixture of the colours of the individual filters as is the case with additive mixtures. In a trichromatic colorimeter, for example, all of the light transmitted by each filter (if a filter type of instrument is used) goes to the matching field and contributes to the colour seen there. The details of the spectral absorption of the individuel filters are immaterial; it is their colours alone which matter. With the subtractive method the colonra of the individual filters are immaterial; it is the nature of their spectral absorption curves that matter. exemple, a certain blue glass and a certain yellow glass in series may transmit green light. This happens because the blue glass transmits the blue portion of the spectrum, a considerable proportion of the green, and none of the red; two stendards in the other beam. The method while the yellow glass transmits no blue, but of reduction of observations recommended by varying proportions of the green to red region, the mokers is based on the following considers yellow filter will give green in combination, as it orango red, orange and yellow-orange, would be possible to have two filters, identical in bimotions of yellow and blue pass through their colours with hose just considered, which in yellow green, green and green-blue, while combination would let through no light at all, combinations of red and blue pass through red. This would be the case if the yellow filter, instead violet, violet and blue-violet Suppose a of transmitting a broad spectral region including the green and red, transmitted only yellow light confined to a narrow range of wave-length. subtracted from all, giving the result as 15 More must be known, therefore, about filters plus 45 red plus 105 yellow. The 45 red is than their individual colours before the colours the nassociated with on equal amount of the of subtractive combinatious can be predicted. Each filter must transmit light to varying extrate over a large region of the spectrum in neutral plus 45 orange plus 60 yellor by order that some parts may be transmitted by all oldsamed. If no the other hand, it is necessary of them. Filters may be found that the to darken the specimen with, say, 003 neutral. necessary properties, and the first successful and that this is matched by 45 red and 25 attempt to utilize this principle for colour him, the 0.03 units of neutral tint are lated measurement was made by 2. W. Loviboud separately as brightness while the colour on the & Sons, London) in the instrument known as the tintomrter. In this instrument the light from the specimen enters one side of the matching brightness adjustment is effected independently field while light from a white diffusing surface, of the Lovibend glasses. This enables the after passing through various coloured glasses in colour match to be made with glasses from not series, enters the other. There are three sets of more than two of the three series and eliminates these glasses. In appearance they may be the use of neutral glasses. This modification

colour has been found by trial the coefficients roughly described as red, yellow and blue, the in the Ostwald equation have to be determined. red having a tinge of blue in it. Each set The methods employed are somewhat com- consists of a series of glasses of graded density, placed and are given in Part II of Ostwale marked with a number proportional to the book." Colour Science." of which an Englah density. That is to say, the glass marked I on translation by J. Scott Taylor has been puls should have the same colour as ten glasses lashed by Messrs. Window & Newton. Equipment of the officerate state of the different sets and the same contract by Ostwala method can, and as on. The numbers of the different sets are so related that a combination of similarly numbered members of each set is of a neutral grey colour. In the original form of the tintometer the glasses were supplied separately and had to be selected and dropped into appropriate slots in the body of the matching also be effected by passing white light through a instrument, which was of a simple form, having number of colour filters in succession. The first filter subtracts some of the spectral constituents distance apart. The full set of glasses for use from the white light; and what is left is robbed with this equipment contained 155 members of each series, graded from 001 to 200, or 465 glasses in all. This degree of subdivision is superfluous, and in the most recent model of the tentometer, known as the British Drug Houses' pattern, 60 red, yellow, and blue glasses, and 12 neutrals are supplied. These are fitted in sliding frames in a matching instrument, which has the comparison fields juxtaposed, giving greater accuracy of matching than the older models In using the tintometer an ottempt is made to match the light from the specimen by a suitable combination of red, yellow and blue standards interposed in the beom from the standard white plate. If the hrightness factor of the specimen is not too great this can be accomplished. If the specimen is too bright to he matched in this way, neutral glasses are inserted in the beam from the specimen, the density being increased until it is possible to effect a match with a combination of The only part of the spectrum which both tions. Any combination of red and yellow glasses transmit is the green, and it is a common glasses alone produces hues intermediate mistake to think that any blue filter and ony between red and yellow, passing through specimen matched by 60 red, 120 yellow and I 5 blue. The smallest of these numbers is first yellow, yielding the resultant 4.5 units of orange. and, as the final designation of the colour, 1.5 neasurement was made by J. W. Lovibond separately as brightness while the colour on the "Measurement of Light and Colour," Geo. Gill nther aide is analysed as 35 violet and 10 blue.

In the latest model of the tintometer the

the measurement. The illuminant provided is the C.I.E. standard B, and the direction of illumination and observation are also those recommended by the Commission. Any measurement made with this instrument can, of course, be converted to the C.I.E. trichromatic system by calculations based on the spectral absorption of the Lovibond glasses. The necessary cal-culations may be made once and for all, and the results incorporated in charts from which corresponding values on the two systems may be read directly. Such charts have now been

prepared by the makers. Many special models of the tintometer for limited purposes are also made. These are supplied with a small number of standard glasses specially adapted for the purpose on hand, such, for instance, as the measurement of X-ray dosage by the Sabouraud-Noire pastille method; the acid washing tests of coal tar fractions; the tests of petroleums and lubricating oils; the evaluation of vitamin-content of codliver oil; or Nessler's ammonia test, and many others. It is probably in these restricted forms, where, owing to the small range of glasses required, the apparatus can be supplied at a much lower price and in more compact form than any universal colorimeter, that the tintometer is most widely used. The degree of reproducibility of different sets of Lovibond glasses, though by no means so good as is sometimes claimed, is nevertheless fairly satisfactory. Any Lovibond measurement can be converted to the C.I.E. trichromatic system by calculations based on the spectral absorption characterising the glasses.

In the Eastman colorimeter, devised by L. A. Jones (J. Opt. Soc. Amer. 1920, 4, 420), the same principle of colour measurement is employed. The graduated series of coloured glasses used in the tintometer are here replaced by three coloured wedges which vary in density from one end to the other. The absorptive properties of the wedges are similar to those of the red, yellow and blue series of Lovibond glasses.

In concluding this brief account of colourmeasuring apparatus and the principles on which they operate, it must be insisted that the successful application of colorimetry to the problems of industry is impossible without adherence to a rigorous system of standardisation as regards conditions of test, and the form in which results are expressed. The degree of standardisation effected by the Commission Internationale de l'Éclairage, though not yet perfect, embodies the pooled experience of the standardising laboratories of the principal industrial nations. Their recommendations constitute a necessary minimum, and should be strictly followed by everyone employing colori-metric methods in industry. New instruments should not be adopted because of ease of manipulation or precision of repetition unless they are designed to work under the standard conditions of the C.I.E., and unless it is possible, by suitable methods of conversion, to express any result obtained with them on the standard reference system.

greatly increases the facility and accuracy of] used for the estimation of substances in solution. The many specific tests of this kind to which colour comparators are applied in chemical and biochemical laboratories do not come within the scope of this article, which is concerned only with instruments and the principles of their use. One of the simplest devices for work of this kind is merely a series of testtubes containing solutions of known and graded concentrations of the substance to be estimated. The solution of unknown concentration is put in a similar test-tube and visually compared with the tubes of known concentration until that nearest to it in colour is found. This method is capable of surprisingly good results if the standard solutions are suitably graded and is still used in many laboratories. Nevertheless, more precise results are to be

expected if an optical comparator is used, and such instruments have been designed. Many of the earlier forms were of crude design, and it is doubtful if they effected an increase in accuracy sufficient to compensate for the extra complexity of technique. Omitting an historical résumé, only typical instruments illustrating the various principles which are employed for chemical estimations by colour or an equivalent criterion at the present time will be described.

Probably the best known instrument is the Duboscq colori-meter, which is based on a principle also used in instruments by Patterson (J.S.C.I. 1890, 9, 36) and Schreiner (J. Amer. Chem. Soc. 1905, 27,

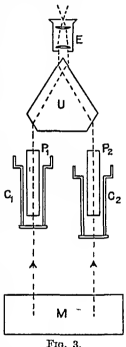


Fig. 3.

1192). Fig. 3 is a diagram of the optical system used in a modern model of the Duboseq instrument. E is an eyepiece, focussed on the sharp edge of a Ulbricht-prism U, which consists of the major portion of a rhomb of clear glass. P₁ and P₂ are "plungers" consisting of glass rods with plane polished ends. These dip into cups C1 and C2, the lower ends of which are plane-parallel glass plates. The plungers are fixed but the cups can be independently moved up and down by rack and pinion adjustments, scales being provided which indicate the distance from the bottom of each cup to the end of the plunger which it surrounds. M is a mirror, inclined to the plane of the diagram, which throws light vertically upwards. The beams passing through the cups Colour Comparators.—Instruments of this and plungers follow the directions indicated class, often misnamed colorimeters, are mainly by the broken lines, and illuminate the two

halves of the field of view, which are sharply from which the result is deduced. Here also, if divided by the top edge of the Ulbricht rhomb. the standard is of different material from the When both cups are empty the light losses in test solution, Beer's law need only be true for the two paths are equal, so if the illumination is the standard and not necessarily for the test properly adjusted the two fields will appear solution, equally bright and the dividing line should be An all nearly invisible. A solution of known cooceu tration is put in one cup, which is adjusted standard solutions except for the initial calibrauntil the length of liquid column traversed by the light has some convenient value. The colour match a brightness match in approxisolution of unknown concentration is put in mately monochromatic light. The colorimeter the other cup, which is racked up or down of van den Bergh and Grotepass is similar in until the two fields exactly match in colour If the unknown concentration is the same as and piniou adjustment moves both cups simul-that of the standard the match will occur when tancously, maintaining both liquid columns of the liquid columns are of equal length. If the concentrations differ the lengths will differ and the nnknown concentration is deduced from that of the standard by assuming Beer's law that for a given colour the product of the concentration and the thickness of the absorbing layer is constant This, the normal method of using the colorimeter, requires the preparation small spectral regions. The depth of the of standard solutions of the test substance. If liquid columns is brought to some predeterthese ere unstable, they must be frequently renewed. To obviate this need more permanent stendards for various colorimetric for which the absorption of the test solution is estimations may be made from coloured glass, or from solutions of some other substance, having the seme colour as a specified con-centration of the test substance. Such standards may differ in their spectral absorption curve found which gives the closest opproximation from that of the test substance, in which to a brightness match. Any residual difference ease the colour match will depend on the in brightness is removed by a small adjustment observer's colour vision end on the spectral of the depth of the liquid columns. It will be quality of the illumination. Difficulties from observed that this method eliminates the the latter couse may be evoided by preparing absorption of the solvent as both beams always and using the standards with an illumination of definite spectral energy distribution, one of the relative sensitivity of these methods depends C.I.E. standards, for example. Standard on the nature of the spectral absorption of the illuminant B will give more sensitive colour discrimination for the majority of estimations gives the more sensitive criterion of concentratban either A or C.

In cases where the test solution does not accurately obey Beer's law, there may be an advantage in using standard solutions of some other substance, even when this is not dictated colorimeter can be used by the second method by other considerations If a standard solution of suitable colour can be prepared from constituents which do obey the law, we can reverse the usual procedure and vary the depth of the standard solution, maintaining a fixed specified

depth for the test solution. The Autenrieth-Konig. Autenrieth-Königsberger colorimeter (Chem. Zentr. 1910, 1, 2032) operates on the same principle as the Duboscq, though the matching arrangements are quite different. Light enters the instrument by two horizontal slits. Behind one of these is a parallel-sided glass cell conteining the test solution, and of purposes including chemical estimations behind the other a long wedge shaped cell either by colour match or monochromatic containing the standard solution. The wedge brightness match. Its optical system is illustrated to the containing the standard solution. cell can be moved past the slit by a rack and trated in Fig. 4. Two beams of light, A and A', pinion adjustment, a scale indicating the thick- enter the instrument ria the lenses Li, Li pinon adjustment, a scale inducating the third.

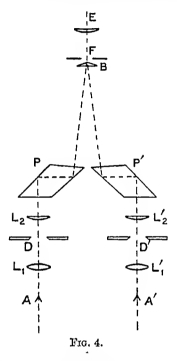
ness of liquid inversed by the beam. The two and then pass through disphragms D. U.
beams ere brought together by a sustable These disphragms constitute the essential
optical system to illumnate the two balves measuring part of the instrument. The aperof the matching field. Beer's law, applied to
turn are square and can be varied in measuring the remaining center on the two optic
whole still remaining center of the two optic

An alternative method of visual estimation. which has the advantage of requiring no tion of the apparatus, employs instead of a design to the Duboseq except that a single rack equal depth One cup is filled with the test solution and the other with the pure solvent A rotatable disc carrying five uentral filters made from gauze of known transparency is fitted in the path of the beam which traverses the solvent, and above the eyepiece enother disc carries eight colour filters transmitting only mmed value, and the colour filter whose band of transmission is nearest that part of the spectrum greatest is brought over the eyepiece fields now appear of the same colour but differ in brightness. The various gauge filters are inserted in the comparison side until one is traverse the same thickness of liquid. The test substance. In some cases a colour match tion and in others the monochromatic brightness match. It is useful to be able to apply whichever of these methods is more sensitive in any given case, and a Duboseq or Autenneth with the aid of a set of colour filters and a series of neutral filters, either of gauze, as used in the van den Bergh-Grotepass instrument, or of lampblack in gelatine. The exact equality of the hound columns maintained in that instrument, though a useful feature, is not a primary require-ment, and may be achieved sufficiently closely by simultaneous independent adjustment of the

cups of a Duboseq type of instrument.

The Pulfrich photometer (" Licht und Lampe,"
1925, No. 3) or Stupho photometer, as it is alternatively termed, can be used for a wide variety between colour, concentration, and thickness axes, by means of micrometer sereus provided

read the areas of the apertures directly. These areas determine the amounts of light which the apertures transmit. This light is collected by the field lenses L₂, L₂', and the two beams are brought together by the internally reflecting prisms P, P', to provide uniform illumination of the two sides of the matching field formed by the biprism B, the ridge of which is focused by the eyepiece E and constitutes the dividing line. A rotatable disc (not shown) contains



eight spectral colour filters any one of which, or a clear blank, may be brought into positions

over the aperture F.

The instrument can be used either horizontally or vertically. In the latter case the light beams are directed upwards by an inclined mirror fitted to the base. For chemical estimations by colour match special cells are supplied. These consist of tubular glass walls fused to optically plano base-plates and fitted with dipping covers which give definite fluid thicknesses of 1, 10, and 50 mm., and a cell in which the thickness can be varied by a solid glass plunger which can be made to descend to various depths by rotation of the cover. The test solution is put in this cell and the standard solution in one of the invariable cells. The two fields of the photometer arc made to match in intensity for white light. This can be done either by independent adjustment of the drums or by setting the drums to equal readings and adjusting the position of the illuminant until both fields are equally bright. The cells are then inserted in the beams (a suitable stage is provided) and the colour match obtained by varying the depth of the test solution as absorptiometer "which can be used at any

with large drums outside the box graduated to with the Duboscq colorimeter. For the monochromatic brightness method two similar cells of equal depth, respectively containing the test solution and the solvent, are used. The most suitable monochromatic filter is inserted, and the intensity of the brighter field is reduced by operating the appropriate drum until a brightness match is obtained.

Accessory attachments for various other types of measurement can be obtained for the Pulfrich photometer. Reference has been made earlier to its use in conjunction with the Ostwald colour standards. Description of these other functions of the instrument would, however,

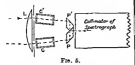
be outside the scope of this article.

monochromatic brightness matching method of estimation clearly reduces to a determination of the ratio of the transmissions of a specified thickness of the test solution and of the pure solvent for approximately monocbromatic light within a selected wavelength range determined by the filter used. The method lends itself readily to accomplishment by purely physical means. An extremely simple method is to employ a photoelectric-cell, of the rectifier type commonly used as photographic exposure meters, and a quickresponse galvanometer having a steady and reproducible zero. Cells containing the solution under test and the solvent are placed in succession in the path of a beam of light from a small gas-filled lamp, such as a 50-watt motor car headlight lamp. The appropriate member of a set of spectrum filters is also inserted in the beam. The relative transmissions of the test solution and solvent for this light may be deduced from the ratio of the galvanometer deflections obtained. If care is taken to arrange the optical system so that the beam passing through the liquids is approximately collimated, and is not obstructed or scattered by the walls of the liquid cells, and so that the beam falling on the surfaces of the receiver is not displaced on interchanging the cells, very satisfactory results can be obtained.

Photo-cells of the kind mentioned require no batteries or auxiliary electrical equipment. Various types differ greatly in spectral response. For this purpose those with selenium surfaces are best, as they have a fairly good response throughout the visible spectrum. Compact self-contained colour comparators employing rectifier photo-cells are now sold by several British makers; see article on Optical Instruments shown at the twenty-eighth annual exhibition of the Physical Society in January 1938 (J. Scientific Instruments, 1938, 15,

For tests of solutions the maximum absorption of which is outside the visible spectrum a rectifier cell is unsuitable. If the absorption is in the infra-red, a thermopile may be substituted for the photo-cell, and, for the ultraviolet, a photo-cell of the emission type, but a difficulty arises in the great scarcity of filters suitable for isolating narrow spectral regions in either the ultra-violet or infra-red spectra.

solvent. On emerging from the cells the beams | plementary to those which are absorbed. are deflected to the sht of a spectrograph hy the quartz rhomboidal prisms P. P', which are mounted one above the other. This slit is of special design, consisting of aeven short alits in



line. One of the beams-that through the solvent-illuminates a short six about 05 mm. square. The other beam illuminates a series of six slits, each 0 5 mm. long, separated by 1 mm, and varying in width from about 0.5 mm, downwards, the width being halved at each step. Seven narrow spectra are therefore ohtained at one exposure. One of these is of the light transmitted by the test solution, the other six, which vary in density in accordance with the widths of the alits, is of the light transmitted by the solvent If at a particular wave length the photographic density in the test spectrum is found to he the same as that in the comparison spectrum from a slit the width of which is only a quarter of that of the test sht, it can be deduced that the illumination per unit area of the comparison beam is four times as great as that of the test beam, and that the relative transmission of the solution to that of the solvent is, therefore, 25%. The use of a microphotometer for determination of plate densities is recommended. This not merely increases the accuracy of the companson, but enables interpolation to be performed. For work in the nitra-violet a source giving a continuous ultra-violet spectrum. such as a high intensity hydrogen lamp, must be employed. The phetographic method is not nearly so convenient as visual methods, or absorption hand in the red, yellow, and green-a direct reading physical methods, but is practiced colour by one showing absorption in the tically the only one which can be satisfactorily blue end of the spectrum. used in the ultra-violet region of the spectrum.

COLOUR AND CHEMICAL CON-STITUTION. produced by the action on the retina of the eye of some portion of white light which has been

part of the photographable apectrum. The octave. White light need not be altered in essential parts are illustrated in Fig. 5. A any way by reflection or transmission from or aboutce giving a continuous spectrum in the through a medium and may thereby be quite. required spectral region is mounted about 60 cm. unaffected. On the other band, certain arefrom the quartz lens L. The light transmitted gularities of surface or certain properties of a hy the lens is restricted by disphragms to two solution may lead to the absorption of a portion circular beams of smaller diameter than the of white light causing the rays refracted or cells C, C', which hold the test solution and transmitted to have colours which aco com-

Such colour, which may be termed " physical colour" because it is produced by physical means and is entirely independent of chemical atructure, is shown by a variety of natural substances, such as hlue eyes, the coloured feathers of a hird, the brilliant green reflex of the wing cases of certain beetles, the colour of butterfiles' wings, and of the rambow. This type of colour can always be recognised, for example, in a hird's feather, by the fact that the colour is only produced by diffraction, and disappears when the object is viewed by transmitted light. There is, however, a type of colour which may be regarded as chemical in origin, that is to say, it is produced by intramolecular interference such as that already mentioned. It is, therefore, dependent on molecular structure, that is to say, on chemical constitution, and is the type of colour with which this article deals. This kind of colour also occurs naturally as hæmin, the colouring material of blood, as chlerophyl, the colouring matter of green grass and as most of the colours found in coloured flowers. These last can be extracted from the object in which they occur, usnally as glycosides, in which the pigment is present in combination with one or more molecules of a monesacchande.

It was clearly understood that the distinction between the different kinds of celour mentioned in this article is made marely as a matter of convenience; there is, of course, no real difference between them except in the method of production. Moreover, there is no sharp division in the matter of their natural occurrence, because the feathers of certain parrots have been found to contain extractable pigments. A chemical substance appears coloured when it gives definite absorption hands in the visible region of the spectrum, the colour being compounded of the colours which remain after those represented by the absorption bands have been removed. A blue colour is given, therefore, hy a substance which gives an

It is ohymus, however, that phenomenon of this kind cannot be confined to the single octave representing the visible spectrum, but must be The sensation of colour is spread over the whole range of the electromagnetic spectrum. Unfortunately, we have no means of detecting absorption bands outside of some portion of water agint water and seven means of detecting asserption founds outcome seattered through the sense of a medium by the visible fregion, except in the ultra-voict which the light has been dispersed either by so far as it is assecptible to the photographic or so octaves of the electromagnetic spectrum by which have been mapped, only one, namely, that, and in the infrared so far as they can which have been mapped, only one, namely, that detected by heat, measurements. There which have been mapped, only one, namely, that is detected by heat, measurements. There is no statement of the properties of the between 3,900A° and 7,600A°, represents the outside the visible region, which made they visible spectrum, or, in other words, the retina give nn absorption in this region are apparently is only sensitive to wave-lengths within the colonriess, but on account of the fact that they give absorption outside the visible region are to he regarded as possessing "invisible" colour, or, in other words, colour that cannot he

detected by the eye.

The hydrocarhon henzene (C₆H₆) is a case in point. This substance shows little absorption throughout the visible region of the spectrum, and therefore appears colourless to the eye. If, however, a photograph is taken in the ultra-violet region it is found that definite marked absorption hands occur, meaning that if the eye could detect waves in this region henzene would he strongly coloured.

It is assumed that this absorption is caused by the rhythmic interchange between the two "Kekulé" individuals I and II, which repre-

sent the formula of henzene, and accounts for its symmetrical properties as well as its stability. It is common to all derivatives in which this interchange is possible. If, however, the interchange is restricted as, for example, when henzene is oxidised to quinone, the ultra violet absorption is influenced, and the absorption thrown into the visible region of the spectrum thus producing colour. Taking, therefore, two comparable examples in hydroquinone and quinone, there is in the one the rhythmic vibration due to the henzene structure, and in the other the absence of such vibration and the occurrence of colour.

Rhythmic vibration producing Rhythmic vibration producing absorption but no visible colour.

Rhythmic vibration producing absorption in the visible region of the spectrum.

Still more striking examples of this change are shown by phenolphthalein:

"Colourless" crystalline substance showing marked ultra-violet absorption.

Deep red substance showing marked absorption in the visible region but diminished ultra-violet absorption.

and also by the colouring pigments of certain flowers, which may he red or blue depending on the formula of the anthocyanidin present:

It follows, therefore, that absorption in the visible region of the spectrum, and the occurrence of visible colour, are mainly dependent on the degradation of certain forms of rhythmic vibration, and if this is true it follows that aromatic substances, that is to say, the derivatives of henzene and allied hydrocarhons in which rhythmic vibrational movement can occur, are the only substances from which compounds having visible colour can be obtained. But this is not the case, hecause many aliphatic substances are also visibly colonred, as, for example, diacetyl, CH₃·CO·CO·CH₃, glyoxal, CHO—CHO, and so forth. An examination of such substances shows, however, that they are all compounds having a conjugated system of double linkages, and on this account are allied to benzene:

It is possible, therefore, that some kind of rhythmic vihration, such as that which gives rise to the ultra-violet absorption of benzene, is also present in these cases, for example:

but that the interchange, being too feeble to colour he termed chromophores, and molecules lead to ultra-violet obsorption, shows its containing them were colled chromogens, Ha influence in the visible region only. Perhaps specified the following chromophores and gave this explanation supplies a reason why certain oubstances which should contain the conjugated chain are colourless, for example, ethyl oxalate (I); the reason being that there is notendency in this case for the substance to assume the tautomene form (II), and rhythmic vibrations are therefore absent.

It is not proposed to deal in this article with the case of occurrence of colour in inorganic compounds The molecular condition of these substances is so complicated that it is not possible in our present state of knowledge to envisage whot may happen when a beam of whits light passes through, or is reflected from, an inorganie erystal in such a way that the emerging beam is coloured Usually the cause of colour may be ascribed to the presence of a coloured ion, such as, for example, ehromum or copper, but even the molecular condition of the salts of these elements vames aufficiently to cause colourless substances to be produced from them by simple reactions. Thus, although copper sulphate is blue under normal conditions, it loses all visible colour when the fiva molecules of water of erystallisation are removed by heat.

GENERAL OUTLINE .- The first synthetic dya (mauve) was produced by Sir W. H. Perkin in 1856. This branch of synthetic chemistry developed with great rapidity, and in the next twenty years the first members of many important groups of dyes had been prepared, e.g. fuchsin (1856), rosanilm blue (1860), and alkylated triphenylmethane dyes (1861-1866); Bismarck brown (1863), chrysoidine (1875), and naphthol orenge (1876) in the aze-group; Martius yellow (1864) in the nitro-group; Magdala red (1868) in the azine series; alizarin (1869) of the anthraquinone group; fluorescella and eosin (1871) representing the phthaleins; and Lauth's violet (1876), foreshadowing the thiazines. By this time sufficient material had been accumulated to allow the formulation of the well known "Witt's rule" (1876), secording to which substances attained the potentiality of colour when their molecules contained certain atomic arrangements or groups, which Witt termed chromophores. The colour and dyeing properties were made manifest by the introduction of additional groups termed suzo- stood.

two things are required to confer on an to bring out the colour and dyeing properties so as to contain this grouping. The quinonoid The atomic groupings giving the potentiality of grouping may be defined as a benzene nucleus

illustrations of chromogens containing them;

Chromosphers
NO 3.

$$-N=N$$
 $-N=N$
 $-N=N$

It will be noticed that some of the chromogens are colourless, e.g. nitrohenzens and phenolphthalem, whilst others have a feeble colour. eg azo-benzene and anthraquinone; none of the ehromogens has any dyeing property. The colour is developed or strengthened by the introduction of salt-forming radicels into the ehromogens. These radicals he therefore termed auzochromes. Hydroxyl, amino, and substatuted amino-groups can act as auxochromes, while sulphonic and carboxyl groups are without this function The following table shows how the chromogens already mentioned era converted into dyes by the introduction of auxochromes:

Witt's Rule is an admirable "working hypothesis" hut it deals only with the connection between a coloured aubstance and its eapacity for affixing itself to the textile fibres and leaves all questions concerning the cause of colour in the chromophore unanswered. Nor, indeed, is the "vibration hypothesis" mentioned above anything more than a reference to a physical condition which is not clearly under-

THE QUINGNOID AND MODIFIED QUINONOID Witt pointed out (Ber. 1876, 9, 522) that THEORIES .- In 1888 (Proc. Chem. Soc. 1883, 27-33) H. E. Armstrong pointed out that the organic substance colour and dyeing properties. chemical formulæ of the well-known dyestuffs The molecule must possess a certain grouping of atoms to give it the potentiality for colour could be elightly modified or restranged, and it must also possess a sail-forming radical accordance with accepted ideas on toutomersm. double bonds in the para- or ortho- position:

Armstrong was probably much impressed by the then recently discovered fact that benzene-azo-β-naphthol was identical with the phenyl-hydrazone of β -naphthaquinone, so that there was as much justification for the

as for the more usual one based on the formation of the substance from β -naphthol and a benzenediazonium salt, viz.:

and if it were allowable to assume tautomerism to a quinonoid form in this case, it might be allowable in other cases, e.g. in the case of p-rosaniline it would be consistent to suppose that on the conversion of the base into salts there was elimination of a molecule of water and formation of a substance with a quinonoid structure:

$$\begin{array}{c} \text{HO}\text{--}\text{C} \overset{C_6 H_4 \cdot \text{NH}_2}{\leftarrow} \\ C_6 H_4 \cdot \text{NH}_2 \\ C_6 H_4 \cdot \text{NH}_2 \\ \xrightarrow{\text{HCI}} C \overset{C_6 H_4 = \text{NH} \cdot \text{HCI}}{\leftarrow} \\ C_6 H_4 \cdot \text{NH}_2 \\ \xrightarrow{\text{HC}} C_6 H_4 \cdot \text{NH}_2 \\ \end{array}$$

In many cases the ordinary formula already contained the quinonoid structure.

The theory was the first real contribution to the relation hetween colour and constitution, because it indicated that the formation of the quinone structure was one of the influences which affected the rhythmic vibration of the Kekulé individuals of benzene so as to throw the absorption from the ultra-violet into the visible region of the spectrum. It is immaterial whether this process leads to visible colour or not, because in certain cases—a fact which has been used to disprove the theory—derivatives possessing undoubted quinone structure are colourless, meaning, of course, that in these cases the "shift" of the absorption from the ultraviolet has not been sufficient. Moreover, opponents of the "theory" have used the argument that it cannot be correct because there are coloured substances, such as azobenzene, which do not possess a quinone structure. This criticism could, with equal force, be applied to copper sulphate. Obviously the formation of a quinone ring from a substance already containing "potential colour" may

to which other atoms or groups are attached by | not. It was never claimed that all coloured organic compounds possessed a quinone ring, although, as a matter of fact, such a statement would not be far removed from the truth.

The literature contains many other hypotheses, which are not theories because the exceptions to them are numerous. It is evident that one cannot distinguish between substances showing absorption in the visible region and those giving absorption in the ultra violet and infra red region of the electromagnetic spectrum. The only truly colourless substance is one showing general absorption throughout the 68 or so octaves of the electromagnetic spectrum. Disturbance leading to absorption may occur in any octave although unfortunately our instruments are not sufficiently sensitive to map more than a small part of the spectrum. This is, however, a physical question and will be answered in time

by the physicists. COLUMBITE. A columbate (niobate) of iron, usually containing some manganese (" manganocolumbite") and tantalum, general formula being (Fe,Mn)(Nb,Ta)2O6. With an increase in the amount of tantalum, there is a transition to the isomorphous species, tantalite. The orthorhombic crystals are dark brown or black and opaque, and range in sp.gr. from 5.3 (columbite) to 7.3 (tantalite). It was in this mineral that C. Hatchett discovered, in 1802, the element columbium, so named because the specimen he examined came from America. Columbite is usually found as single crystals embedded in pegmatite or granite; for example, at Haddam and Middletown in Connecticut, Chesterfield in Massachusetts, brilliant crystals in granite at Standish in Maine, large masses up to 2000 lb. in granite veins in the Black Hills of South Dakota, and at several other American localities. At Ivigtut in Greenland it has been found in the cryolite deposit. Large crystals are found in the felspar quarries near Moss in Norway. Other localities are Rabenstein in Bavaria, Bushman Land in South Africa, Madagascar, etc.

L. J. S. COLUMBIUM, NIOBIUM (Sym. Cb, Nb). At. wt. 92.91. The name columbium is used in current American literature, although the element is now known as niobium in Great Britain, France, and Germany. The first name is due to Hatchett, who in 1802 isolated a new "earth" (Nb₂O₅) from a mineral found in Connecticut. He called the mineral columbite and the element columbium. It was not until 1844 that Rose (Pogg. Ann. 1844, 63, 307, 693) observed a wide variation in the density of many tantalites and columbites, and of the acids derived from them, and realised that they contained two closely related elements. It was Rose who introduced the name niobium for Hatchett's columbium. In all probability Hatchett's "earth" was a mixture of Nb_2O_5 and Ta_2O_5 . The name niobium and the symbol Nb will he used throughout this article, in accordance with the nomenclature adopted in the First Report of the International Union of Chemistry, 1936.

Niohium is a member of a group of comlead to the production of visible colour or may paratively rare metals, including tantalum, tound in the oldest Flution locks, and especially which contained some 3% of aluminium and in and around intrusions which traverse permatities and other extremely coarse-grand caroum. This treatment suffect to vaporase granito rocks. Monazite, and other minerals the impurities out of the nichium. The metal containing metals of the cerium group, zircon, eryolite, and fluorspar, and minerals continuing K,NbF, with hydrogen. The product is a lithium, easuum, and runhulum, aro often black powder from which residual hydrogen associated with these minerals, and are probably may be expelled by shaping it into small intimately connected with their genesis.

Although a rare metal, niobium exists as the pentoxide in a considerable number of 1927) have described the reduction of Nb.O. minerals, in some of which—such as pyrochlore by calcium in presence of a small quantity of and columnite—it may be regarded as the alkali metal and of calcium chloride. This essential constituent, while in othere, although operation is carried out in a sealed tube at practically always present, its proportion varies from a mere trace up to an amount which justifies its heing considered so the principal oxide.

Nichium occurs in minerals only as the pentoxide, Nb₂O₃, in combination with a base, the mobium in a few minerals (such as wohlerite) which contain silica being present as a silicomobs to and never as a silicate of mobium.

A list of the principal minerals containing niobium will be given in the article Tantalum, with which metal mobium invariably occurs in nature. From the technical point of view the principal michium minerals are those which consist essentially of niobates of iron, manganese and lime, zirconia and metals of the cerium group being also usually present in, or at least intimately associated with, such minerals as contain limo

The mineral richest in nichium is pyrochlore, a mobate of lime containing titanium, cerum, and thorum, together with other alkalmo satisfactory for the electro analysis of niebu cartbs and alkal bases and soom fluorent tantalum mixtures. The niebum depo Tho most important mineral, bowever, is the were resistant to corrosion by mineral acids. niobate of iron and manganese known as columbite or niobite (q v), in which the Nb Os may even exceed 78%.

New deposits are constantly being discovered, and mobium, like many others of the rare metals for which a demand now exists, will doubtless be found to occur in much larger quantities than is now generally known. One property in Weatern Australia is at present being operated exclusively for the production of tantalite nichite concentrates. Considerable amounts are also available as hy-products from the preparation of the rare earths employed for incandescent mantles, from tin and wolfram ores, and from the heavy waste material ohtsined in the dressing of eryolite.

Metallic niobium may be prepared by passing a mixture of the vapour of the pentachloride and hydrogen through a red-hot tube (Roscoe) Moissan reduced the pentoxide by means of fusing the ore with an alkali or alkaline salt, carbon in an electric furnace, but the resulting nichum contained 25-34% of carbon. Von Bolton introduced a method of manufacturing hydrolysis, with production of an insoluble mobium for metal filament lamps in which matters of nichle and tentale acids. The the pentoxide was mixed with paraffin and mixture on ignition gives the pentoxides Nb,O, formed into threads, which were bested in record and Ta₂O₂. The preparation entails the to a temperature of over 1,600° by passage of an irrend in turn of tin, antimony, iron, men. electric current. The use of niobium in lamp ganeso and similar elements, and also of manufacture has now been discontinued.

tungsten, molyhdenum, titaninm, and uranium, by von Bolton hy reducing the pentoxido with commonly occurring together in minerals aluminium powder, and heating the product, is also obtained by reduction of the fluoride cylinders and melting in excuo in an electric furnace. Marden and Rich (U.S.P. 1728941,

900-1,000°. The electrolytic production of nichium from a mixture of nichium and tantalum pentexides was described by Isgarischev and Prede (Z Elektrochem 1933, 39, 283-288). The oxide mixture was fused with potassium pyrosniphate and the melt extracted first with water and then with 20% potassium hydroxide solution. Electrolysis of the alkali extract with the addition of from 1-1 5% of dextrin, with a platinum anodo and a copper cathode, yielded a deposit of pure nichium Alteroatively, the melt was extracted with saturated ammonium oxalate solution, with the addition of a smaller amount of esturated oxahe acid solution, or with citric acid solution, and electrolysed at a temperature above 80° Niobium was thus deposited quantitatively, and tantalum pentoxido could bo recovered from tho electrolyte by acidifying with sulphuric acid. Electrolysis in acid solutions was found to he satisfactory for the electro analysis of nichiumtantalum mixtures Tho niobium doposits

Petrce (J Amer. Chem. Soc 1931, 53, 2810) found that mobium, but not tantalum, could be electro deposited at 103° from a saturated sodium carbonate solution in which the ignited pentoxide had been dissolved. A copper eathodo and a current density of 3 smps per aq dm. were employed. Balko (Ind Eng Chem. 1935, 27, 1168) states that metallic mohium may be produced by minor variations of the electrolytic method used for tantalum The fessed double fluoride KaNbF, is electro-lysed, using a cast iron pot as cathode and a rod of graphite as anode, and adding mohium pentoxide to the molten bath.

The separation of michinm from tantalum is a problem chiefly of importance in con-nection with the preparation of tantalum, and will be dealt with fully under TANTALUM. The general method of extraction consists in extracting the mixture of nichates and tantalates titanium.

Larger quantities of mobium were produced. The separation of nichium and tantalum is

a matter of considerable difficulty, owing to the close similarity of these two elements. One of the commonest methods is that developed hy Marignac in 1866, depending on the different soluhilities of potassium niobium oxyfluoride, K₂NbOF₅, H₂O (1 part in 12-13 parts of water at 17°-21°), and potassium tantalum fluoride, K2TaF7 (1 part in 150-160 parts of water containing a little hydrofluoric acid, at the same temperature). These two compounds are not isomorphous and do not form mixed crystals or solid solutions. In carrying out the separation the mixture of niohic and tantalic acids is dissolved in concentrated by drofluoric acid and the correct amount of potassium fluoride is added. On concentration, the potassium tantalum fluoride separates first and is filtered off. The filtrate is concentrated further with the addition of more hydrofluoric acid and potassium fluoride, and yields crystals of potassium niohium oxyfluoride mixed with potassium tantalum fluoride. These two salts are then separated hy recrystallisation.

Numerous alternative methods of separation have been suggested and applied. Thus, for example, a solution of potassium hexa-tantalate and-niohate is formed by dissolving 1 part of a mixture of Ta₂O₅ and Nb₂O₅ in 6-8 parts of water with the addition of 3-4 parts of potassium hydroxide. From the resulting solution pure tantalic acid is precipitated at 15-30° by the action of carbon dioxide or a hicarbonate (U.S.P. 1908473, 1934). Another utilises the solubility of niobium pentoxide, and the insolubility of tantalum pentoxide, in a mixture of equal volumes of selenium oxychloride and concentrated sulpburic acid (Lehner, J. Amer. Chem. Soc. 1921, 43, 21; Merrill, ibid. 1921, 43, 2378). Hydrolysis of the extract after dilution yields niohic acid. Ruff and Thomas (Z. anorg. Chem. 1926, 156, 213) found that niobium pentoxide is converted to the pentachloride by beating at 200-225° with carbon tetrachloride, whereas tantalum pentoxide is unaffected. Unfortunately, the metals cannot he separated by this means, as niobium pentachloride reacts with tantalum pentoxide, forming niohium oxychloride and tantalum pentachloride, which are not readily separated.

Metallic niohium melts at 1,950° (von Bolton, Z. Elektrochem, 1907, 13, 145) A lower value (1,700°) is recorded by Guertler and von Pirani, (Z. Metallk. 1920, 12, 67; Ann. Phys. 1915, [4], 48, 1034). Its electrical resistance is 0.1870 ohm for a wire 1 m. long and of 1 mm. diameter (see also Balke, Ind. Eng. Chem. 1929, 21, 1002). It is less malleable or ductile than tantalum, hut the rolled metal has a hardness greater than that of wrought iron and may he welded at a red beat. Balke (Ind. Eng. Chem. 1929, 21, 1002) states that in the annealed condition niohium is more ductile than tantalum. Many of the properties of niobium as known at present were determined with a sample of the metal prepared by von Bolton (l.c.) by reducing the oxide with aluminium. This author gave the density as 12.7, hut more recent values range from 7.4 to 8.56.

heated in air, some nitride heing formed together with the oxide, hut the rolled or cast metal is hut little affected, as a protective coating of oxide rapidly forms. It is practically unaffected hy any acid except hydrofluoric acid, or hy solutions of the alkalis, but is rapidly attacked hy fused alkalis or alkaline nitrates.

Metallic niohium comhines with hydrogen at a high temperature, and according to von Bolton a hydride, NbH, is formed. Hagen and Sieverts (Z. anorg. Chem. 1930, 185, 225) have since shown, however, that the amount of hydrogen taken up is a function of temperature and pressure, and the formation of a true compound is very doubtful. The compounds of niobium have not yet heen applied commercially, but the pentoxide and the fluorides, especially the double fluoride K₂NbF₇, are of interest in connection with the preparation of the pure compounds or metal from minerals. Niohium in amounts up to about 2% inhihits intergranular deterioration when chromiumnickel steels of the stainless type are exposed to elevated temperatures and chemical corrosion. Tantalum gives similar results, hut a larger proportion of it is required (Becket and Franks, Trans. A.I.M.E. 1934, 113, 143). Addition of niohium to steels containing 4-6% of chromium and 0.1-0.2% of carbon is found to inhibit the air-hardening property of such steels. Niobium bas a limited use in valve manufacture.

Alloys of niobium bave hitherto received relatively little attention. Iron and niobium appear to alloy in all proportions, and ferro-niobium may be formed directly by reducing the mixed oxides. Aluminium-niohium alloys have been produced by the Goldschmidt process, and a brittle alloy of chromium and niobium is formed hy fusing green cbromium oxide and niobium together in the electric furnace (Goldschmidt and Vautin, J.S.C.I. 1898, 17, 543; Moissan, Bull. Soc. cbim. 1902, [iii], 27, 431). Alloys of nickel and zirconium with niobium and tantalum have been prepared, and it is claimed that the alloy containing zirconium can be heated to whiteness in air without oxidation or vaporisation (Canada P. 209342, 214118, 1921; U.S.P. 1334089, 1920).

The removal of tin from Fe-Nb-Ta alloys ohtained in the reduction of niohite is carried out hy fusion with sufficient silicon to convert the niohium and tantalum to the silicide (Nb, Ta)₂Si. The resulting hrittle alloy is powdered and hoiled with hydrocbloric acid to extract the tin. The purified powder may he melted and hlown to remove silicon, or may he used direcly for the treatment of stainless steels 450857, 1935). Ferrosilicon-niohium (B.P. alloys are produced from a mixture of niobite, silicon, lime, and silica by fusion in an electric furnace (B.P. 434400, 1935).

COMPOUNDS OF NIOBIUM.

Niobium monoxide, NbO, has heen described by various authors, but its existence is somewhat ange from 7.4 to 8.56.

The powdered metal oxidises rapidly when K_2NbOF_5 with sodium in an iron crucible, or by reducing NhOCI, with magnesimm, and is base, and adding alcohol. They are white described as a black material. It burns to the powders stable in contact with air, and decompentoxide when heated in air, ignites in chlorine, posed by water. forming NhOCl, and is soluble in hydrochlorio acid with evolution of hydrogen.

Nioblum sesquioxide, Nb.O., is formed by reducing mobium pentoxide with magnesium

of hydrogen at 1,250°

heated to redness in the air.

Nioblum pentoxide, Nb₂O_g is a white infusible powder of spg r. 453-48, according to the method of preparation. It is formed by direct oxidation of mohium, or of its lower oxides, and also, in the hydricities of pentavalient inohum saits. The molecular heat of formation inohum saits. The molecular heat of formation from the elements is 441,330 g -cals strong ignition mobium pentoxide is insoluble in all acids except hydrofluoric acid. It dissolves, bowever, in molten potassium bydrogen aulphate, or ammonium hydrogen sulphate, and in fused alkali hydroxides or carbonates, although it is insoluble in solutions of alkalis The mobium pentoxide content of a mixture of Nb₂O₃ and Ta₂O₃ may be determined by reducing the Nb₂O₄ to Nb₂O₃ by means of hydrogen, and determining the increase in weight on ignition. Ta, O, is not reduced by hydrogen

Niohic acid,-This term is applied to the more or less hydrated pentoxide formed by the hydrolysis of compounds such as mohium pentachloride or oxychloride. A sımılar material is formed by the action of mineral acids on alkali monates. The gel is completely dehydrated by heating to 300° (cf. Suc. Compt. rend. 1932, 194, 1745) Freshly precipitated niobic acid gels are soluble in acids and also in

sodium or potassium hydroxide solution.

Numerous mobates have been described, only those of the alkali metals being adulte in water. Sodium metanichate, NaNbO3, is obtained in the anhydrous form hy fusion of mobinm pentoxide with acdium carbonate and acdinm fluoride It crystallises with seven molecules of water of crystallisation. Sunder and Trey (Z. anorg. Chem 1931, 196, 321) have obtained products such BaO, Nb,O, 8.5 4BaO, Nb,O, by heating barium carbonate with mobium pentoxids to 400-500°. Many other niobates have been described (see "Text Book of Inorganie Chemistry," edited by J. Newton Friend, vol VI, part 111, pp. 156-166), hut their compositions and constitutions are in some cases uncertain

Perniohic acid, HNbO, 7H, O, is obtained is obtained. The hydroxide is soluble in excess

Niobium pentafluoride, NbFs, forms colourless, strongly refractive, monochine prisms, d13 3 293, m.p. 75 5°, bp. 217-220°/760 mm, and is obtained by the action of fluorins on powder, or by heating the pentoxide in a stream mobium, or by treating mobium pentachloride with anhydrous hydrogen fluoride in a freezing Miobium dioxide, NiO₂, a black powder, as maxture, and purifying by redustilation (Ruef and Dedner, Ber. 1909, 24, 493; Ruff and Bydrogen, or with magnesium. It huras when Schiller, Z. anorg, Chero 1911, 72, 229), 11 a the solution yielding mobic acid on addition of amroonia or sodium hydroxide. It has a strong tendency to form stable double fluorides with the fluorides of other metals,

Niobium oxyfluoride, NbOF, is obtained by the action of hydrogen chlorids on a fused mixture of niohium pentoxids and calcium fluorade at a red heat. It forms double salts with other metallic fluorides (a.g. Na, NbOF a

(NH4)2NbOFs, ZnNbOFs,6H2O), Niobium trichloride, NbCl, is a non-volatile, non-deliquescent, crystalline substance resembling sodine, and is formed by leading the vapour of the pentachloride through a heated tube It forms long dichroic needles, which are exidused on exposure to air and which yield mohium exychloride, NhOCI,, and carbon monoxide when heated in carbon dioxide

(Roscoe, Chem. News, 1878, 37, 26),

Nichtum pentachloride, NhCls, is formed as yellow needles by heating mobium pentoxide mixed with charcoal in a stream of chlorine, It is also formed by heating the pentoxide with sulphur monochloride in a sealed tube at 200°, and fractionally distilling the products in a current of chlorine. It is produced, together with the oxychloride, by heating the pentoxide with carbon tetrachloride. It has mp. 194 h p. 240 5°, sp gr. 2 73-2 77, and gives a normal vspour density. It is soluble in carboo tetrichloride, chloroform and alcohol. On heating with 3% sodium amalgam in absence of air and extracting with water, chloroniobium chlorode, (Nb₂Cl₁₂)Cl₂,7H₂O, is obtained as black ahming crystals, forming an olive green powder, insoluble in cold water, but soluble in het water to an three green whation. Only two of the chlorine atoms are some. When treated with the equivalent amount of aodium bydroxide, chloroniobium hydroxide,

(Nb,C!,s)(OH)2,8H2O,

is formed as a black microcrystalline precipitate, and from this the bromide,

(Nb,C1,,)Br,7H,O,

as a yellow amorphons powder on decomposing of sodium hydroxide, and from the solution potassium pernichate with dilute sulphure excess of concentrated hydrochloric acid preand and dayling the groduct, or by the cipitates a hrown powder, Nb. Ch., 90 H.O action of 30% bydrogen peroxade on niobue (Hanned, J. Amer, Chem. Soc. 1913, 25, 1078).

and. It loses oxygen at 100°, and forms Reduction of a solution of niobum pentihydrogen peroxids on heating with dilute chicaide at a platinum cathode gives a blue sulphune acid. Perniobates of the alkali colution, which is of a colloidal character; with roctals of the general formula M,NhOa are gold obluride it gives a magnificent purple formed by treating the corresponding niobate liquid, from which, on boiling, a purple prewith hydrogen peroxids and a solution of the cipitate deposits. This precipitate is similar

Niobium pentachloride forms an addition compound with piperidine, NbCl₅,6C₅H₁₁N, and gives similar compounds with related organic bases. It reacts with boiling benzene to give the compounds $3NbCl_5, C_6H_6$, $2NbCl_5, C_6H_6$, and $NbCl_5, C_6H_6$, and $NbCl_5, C_6H_6$, and $NbCl_4(C_{10}H_7)$ and $NbCl_4(C_{10}H_7)_2$. With phenol in carbon disulphide solution the compound

Nb(OC₆H₅)₄CI,

m.p. 233-235°, is formed (Funk and Niederländer, Ber. 1928, 61, [B], 249, 1385). Niobium Niobium pentachloride reacts with glacial acetic acid to form the compound Nb₂O₃CI(OAc)₃ (Funk and Niederländer, Eer. 1929, 62, [B], 1688).

Niobium oxychloride, NbOCl₃, is prepared by the action of chlorine on a mixture of niobium pentoxide and carbon, and forms white silky crystals which can be sublimed.

Niobium oxybromide, NbOBr₃, pentabromide, NbBr₅, also exist, but the formation of an iodide appears to be doubtful (cf. Barr, J. Amer. Chem. Soc. 1908, 30, 1668).

Niobium nitride, NbN, is prepared by heating the pentoxide mixed with the calculated weight of carbon in nitrogen at 1,250°. It is a light grey powder, m.p. 2,050°, which is decomposed by alkali hydroxide (Friederich and Sittig, Z. anorg. Chem. 1925, 143, 293-320). Triniobium pentanitride, Nb₃N₅, is prepared by prolonged heating of powdered niobium in nitrogen at 1,000°.

Niobium carbide, NbC, has been produced by heating niobium sesquioxide mixed with carbon in hydrogen at 1,200°. The molten metal also absorbs graphite to yield carbides of unknown composition. They are brittle and hard, and will scratch glass or quartz (Moissan, Bull. Soc. chim. 1902, [iii], 27, 431). The use of a niobium carbide and a binder in a metal composition has been described (B.P.)

391933, 1933).

Niobium boride, NbB₂, is formed as a hard, grey, crystalline material in the electrolysis of fused mixtures of alkali and alkaline earth borates and fluorides with niobic oxide. Any amorphous boron or calcium boride is removed from the product with dilute nitric acid. The crystals resist nitric and hydrochloric acids and aqua regia, but are easily oxidised and are slowly attacked by fused alkalis, and by cold hydrofluoric or sulphuric acid (Andrieux, Compt. rend. 1929, 189, 1279).

Niobium phosphide, NbP, is formed by heating red phosphorus with mctallic niobium at 550° (Heinerth and Biltz, Z. anorg. Chem. 1931, 198, 168). It burns readily when heated in air. Niobium arsenide NbAs1.s, is prepared by prolonged heating of the elements at 600°.

Niobium sulphide, NbS2, has been prepared as a black powder by heating the elements together. Metallic niobium is not attacked by H2S and CS2 on Nb2O5 at 1,000° (Hönigschmid, applications of condensation was in the pre-

to purple of Cassius; it colours glass a violet- | ibid. 1934, 219, 161). Niobium oxysulphide, red. According to Kiehl and Hart the blue NbOS3, is also formed as a black crystalline solution contains tervalent niobium (ibid. 1928, powder by heating the pentoxide in the vapour of carbon disulphide mixed with carbon dioxide.

For methods of analysing niobites and tantalites, and of detecting and estimating niobium, see Chemical Analysis (Vol. II,

p. 602).

The Arc Spectrum of Niobium has been studied by Hildebrand (J. Amer. Chem. Soc. 1908, 30, 1672); Barr (ibid. 1668), Mott (Trans. Amer. Electrochem. Soc. 1917, 31, 372), and Laporte (Z. Physik, 1926, 39, 128). The most persistent lines ("raes ultimes") are given by de Gramont as $\lambda 4058.97$ and $\lambda 4079.73$ Å (Compt. rend. 1920, 171, 1106). H. J. E.

COLZA OIL syn. for RAPE OIL.

COMFREY, Symphytum officinale. common comfrey (Fam. Boraginaceæ), a riverside plant, has been used since Saxon times as external application for wounds. The air-dry root contains 0.6 to 0.8% allantoin to which its healing action has been ascribed.

COMPRAL. A combination of amidopyrine and trichlorethylurethane (Baeyer Products, London). B.P.C.

CONARACHIN v. Arachin.

CONCHAIRAMIDINE Cinchona Alkaloids (this vol., p. 160a).

CONCHAIRAMINE CINCHONA Alkaloms (this vol., p. 160b).

CONCRÈTE v. Building Materials.

CONCUSCONINE v. CINCHONA ALKA-

LOIDS (this vol., p. 160c).
CONDENSERS. In order to convert the vapour of a substance into a liquid, it is necessary to remove a quantity of heat equal to the heat of vaporisation. The apparatus which effects this transfer of heat is called a condenser.

The shapes and forms of condensers are many and varied, but there are two distinct types under which almost all patterns can be classified.

The two types are:

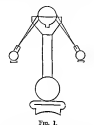
1. Surface condensers, in which the cooling medium is kept separate from the vapour to be condensed, and

2. Jet condensers, in which the cooling medium is supplied directly to the vapour either in the form of a spray or as a thin

It is apparent that the history and development of condensers are parallel to those of distillation methods. It was not until the practice of distillation was improved by the alchemists that methods of condensation received attention. The origins of alchemy can be traced back in the researches of Hoefer, "Histoire de la Chimie," 2nd ed., Paris, 1866; Kopp, "Geschichte der Chemie," 1866; Kopp, Braunschweig, 1847; and Berthelot, "Les Origines de l'Alchimie," Paris, 1885; "Intro-duction à l'Étude de la Chimie des Anciens et du Moyen Age," Paris, 1887; "Collection des Anciens Alchimistes Grecs," Paris, 1887; hydrogen sulphide. A second sulphide, Nb₂S₅, and "Histoire des Sciences—La Chimie au has been reported as formed by the action of Moyen Age," Paris, 1893. One of the earliest

paration of mercury. The preparation of this pillustration of an alembic to which two receivers element is described by Dioscorides. Cinnabar are attached (Fig. 1). was heated in an iron pot and the vapours condensed in an earthenware cover, which was denoted by the Greek word "ambax," The Arabic prefix "al" was added, making the work "alambic," or, as it is known to day, " alembic."

Pliny describes a primitive method which was used for obtaining turpentine from rosin.



Fleeces of wool were hing over kettles, the vaporised oil condensing on the fibres of the wool. Sponges were used by sailors to obtain wool. Sponges were need to sainter to obtain fresh water at sea. Alexander of Aphrodissas (third century), in his commentaries on Aristotle's "Metcorologica," states that, "They boil the sea water and suspend large sponges from the mouth of a brazen yessel to imhibe what is evaporated and, in drawing this off from the sponges, they find it to be sweet water,"



F10. 2.

The earliest aketchea of apparatus date from the time of the Greek alchemists of Alexandria, Zosimns, about the end of the third century, compiled a kind of chemical encyclopædia incorporating the work of previous writers. One of them was a woman alchemist by the name of Cleopatra who lived about the beginning of the Christian era. Cleopatra bad written a of alcohol. Authentic drawings of apparatus treatise on gold-making which was called the of that period have not been found.

"Chrysopea." The "Chrysopeas" contains an luring the sixtentib century changes in the

It is probable that air cooling alone was



Fig. 3.

osed for the straight tube and alembic condensers up to the discovery of alcohol in Italy sometime during the eleventh century. The use of water

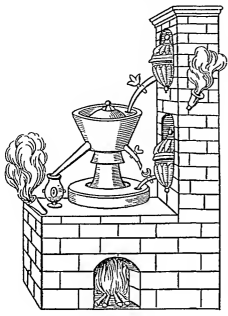


F10. 4.

for cooling purposes was first mentioned by Raymond Lully (1235-1315) in his extensive writings on the curative and clixir properties



simple alemhic condenser design were made to achieve hetter cooling. The first step was to lengthen the delivery tuhe which was then cooled hy passing it through a tuh of water. Fig. 2 is from "Thesaurus Euonymi Philiatri." The next improvement was to pass the tuhe diagonally through the water barrel to increase the cooling area, as in Fig. 3. The earliest illustration of a coil or worm condenser is shown in a work hy Biringuccio (1540), Fig. 4.



Frg. 6.

A similar design is shown in Fig. 5 which is taken from Lonicer (1551).

The first attempt at continuous cooling was tried some time in the middle of the eleventh century. Fig. 6 is taken from the French edition of Matthiolus's commentaries on Dioscorides (1501-1557).

During the eighteenth century there was no substantial advance made in condenser design.

were in general use and were adequate for the uses to which they were put. The counter-current principle hecame hetter known and more widely used during the later part of the nineteenth century. Ettling ("Handwörterhuch der Reinen und Angewandten Chemie," 1842, Bd. 2, p. 531) assumed that Liehig deserved credit for the invention of the counter-current condenser, and to this day the simple straighttuhe condenser is called a Liebig condenser

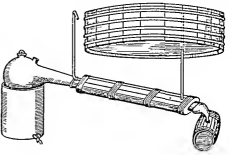
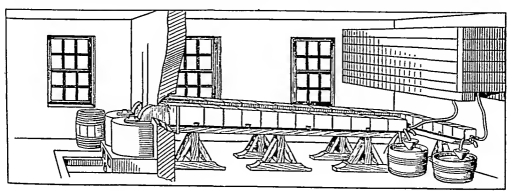


Fig. 7.

(Fig. 9A). Kahlbaum (Ber. 1896, 29, claimed that Weiler was the first to see the advantage of admitting the cooling water at the end of the condenser furthest from the hot vapours. However, Speter (Chem.-Ztg. 1908, 32, 3; Chem. Weekblad, 1931, 28, 381— 382) showed that the principle was first proposed by an unknown French chemist and later developed independently by Weiler and Gadolin. The condenser designed by Gadolin is shown in Fig. 7 (1778). Further investigation by Speter (Chem.-Ztg. 1908, 32, 654) proved Lavoisier to he the author of the anonymous publication cited. Schelenz (Z. angew. Chem. 1910, 23, 1978; Chem.-Ztg. 1909, 33, 141, 154) believes this discovery was made by the French chemist Dariot (1533-1594), but was never appreciated until a later date. Poissonier (1770) also had realised the necessity of providing a large cooling surface and a counter-current flow of cooling water (Fig. 8).

The surface condensers designed since this period do not differ fundamentally in principle The worm cooler and the water-cooled alembic from the ones just described, although the

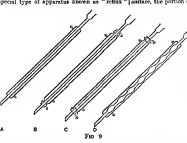


Frg. 8.

differences in detail are vitally important, boiling point (100°C. to 200°C.), it usually The straight condensing tube has been replaced suffices to employ the inner tube merely as an by sprais, bulls, series of bulls (Fig. 2D), com-bunations of these and divers methods of con-bunations of these and divers methods of con-tecting the vapour and the cooling surface, Micros (J. Chem. Educ.) 1930, 7, 2963–29635, over a large area in a minimum of space. A special type of apparatus known as "redia."

cooling jacket is sometimes made of approximately rectangular section. A condenser with expanded sections along the inner tube was described by W. Friese (Pharm, Zentr. 1913, 54, 670). The apparatus consisted of ordinary Liebig jacket mside which was a series of circular chambers with conical tops and bottoms, giving a large cooling surface. condenser for accomplishing the same result is shown in Fig. 9D. The Soublet single bulb condenser which

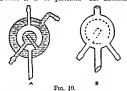
85



condensers was designed to return condensed | reflux condenser is shown in Fig. 10B. An all vapour back to a vessel containing boiling glass spiral pattern is shown in Fig. 11. liquid without any sensible loss of vapour A slightly modified form of Liebig co through evaporation Reflux condensers are extensively used in extraction processes and in organic reactions where the loss of volatile solvent is to be prevented The historical

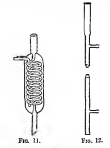
commonly used

A slightly modified form of Liebig condenser is described by West (Ind. Eng. Chem. 1928, 20, 737) The space between the inner and outer tubes is reduced to 1-1-5 mm. and the tubes are scaled together at the ends. ' The water



developments of the return condenser have been outlined by H. Meyer (Chem Ztg. 1910, 34, 351, 421), L. Galates (ibid. 809), and Schelenz (Z. angew. Chem. 1910, 23, 1975-1980). The simplest type of laboratory condenser

that is widely used is the Lieber form shown in Fig. 9A. It consists of a straight cylindrical glass tube in which the vapour is condensed. The upper end is widened out slightly to facilitate connection with the still, and its lower end is cut off obliquely; a cylindrical glass jacket covers the greater portion of the length of the tube and, through it, by appropriate entrance and exit tubes, cold water is circulated in an upward direction. The glass jacket and inner tube may be fused together (9A), or connected by cobber air water cooled one. It was suggested by



consequently passes much more rapidly through the outer jacket and the cooling is found to be more effective (Fig. 12). The apparatus was the inner tube (West, Ind. Eng. Chem. [Anal]. 1930, 2, 199).

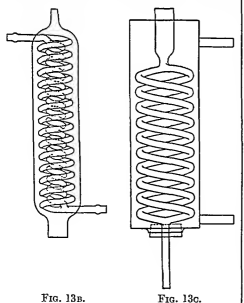
(9B and C). For hauids of moderately high Hopkins (J.S.C.I. 1897, 16, 979) that if the

relative positions of the vapour and cooling stream of water are reversed, a larger area will be exposed. In Fig. 13A the water flows in the direction of the arrows. The outer jacket serves as an air-cooler, while the inner tube is water-cooled. Two forms of condensers are large-capacity laboratory

described by Othmer (Ind. Eng. Chem. [Anal.], 1929, 1, 153) in one of which the cooling is internal by means of a double-coiled tube, with water flowing through both tubes in parallel, thereby giving a maximum amount of cooling surface in minimum space (Fig. In the second type the cooling is external and there is a double-coiled inner tube for the vapour (Fig. 130). Bajda (Ind. Eng. Chem. 1919, 11, 52; the same in U.S.P. 1317262, 1919) passes the vapour through a preliminary air-cooler before passing it through the water chamber.

A number of condensers of the "double- surface " type have been FIG. 13A.

described. The vapours are con-densed in the annular space hetween two surfaces, hoth of which are water-cooled, the design being such that one stream of water flows in succession over each. These condensers can he made quite short; they are readily fixed in a vertical position and occupy little bench space. A modified Soxhlet's hall

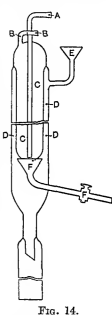


type is shown in Fig. 10A. Several vertical condensers have been described by Friedrichs in which one of the two surfaces between which the vapours condense is constructed spirally (Z. angew. Chem. 1910, 23, 2425-2426; 1912,

from the solvent-extract mixture without changing the apparatus. Bloom (Ind. Eng. Chem. 1910, 2, 103) made such an apparatus (Fig. 14). And more recently Zinzadze (Chem. Fahr. 1930, 113) has designed a doublechambered condenser for extraction or distillation. The top division is cooled hy a coiled water tube and the liquid collects in the hottom of this division and may be withdrawn by a side tuhe, provided with a three-way tap, or returned to the lower division. Various modifications are described in the article (see also Zinzadzé, Biochem. Z. 1930, 220, 177-184, 185-191; Bull. Soc. chim. 1931, [iv], 49, 1204-1205).

The first comparison of the relative efficiencies of laboratory condensers was made by Dover and Marden (Ind. Eng. Chem. 1916, 8, 834-836). The authors reported their conclusions as follows: "The rate of boiling has a very marked influence upon the efficiency of the condenser. The bore of the condenser has

influence upon the efficiency of the condenser as has also a narrowing at the tip or a constriction anywhere the tube. (Noticed in the spiral type where the spiral is fused on to the inner tube.) These conditions tend to cause chocking, and when this occurs loss is always disproportionately great. The length of the condenser is a factor in its efficiency (especially in the case of the Liebig form) for low hoiling-point liquids. In the case of other forms, the length of the jacket has less influence than seems to he commonly supposed. In experiments where a long condenser can conveniently used, the Liehig seems to be preferred because it is a much less costly

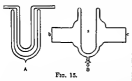


condenser and can be easily cleaned. Where a short condenser is required, the Friedrichs seems best. The short Lichig or Allihn can be used to advantage only when the rate of condensation is not greater than from 2 to 3 drops per second, or when some means is used, such as a glass wool plug in the top of the condenser or a test-tube inverted over the top of the condenser, to prevent too rapid carrying away of the vapour by air currents."

F. Friedrichs (Z. Angew. Chem. 1920, 33, 29-32) presented a critical discussion of the various forms of laboratory condensers which had been described in the literature from the time of Liebig onwards. The screw-shaped condensing tube with five "threads" and provided with internal cooling jacket was most efficient. This form is more suitable for use as a 25, 2200-2209). In certain extraction operal reflux apparatus than is the spiral form. Mach tions it is desirable to remove the solvent and Herrmann (Chem. Fabr. 1931, 4, 157, 170)

type. Results are tabulated for experiments. For this later purpose, a novel form of condenser with nine types of coolers. The best three is described in B.P. 6916/1905, which combines coolers in the order of efficency were: Liebig efficiency with extrems simplicity of con-with four internal tubes, those with bulbs efficiency with Extrems simplicity of con-blown in the internal tube, and those with small paring pure distilled water (so called conjacket diameter. Counter-entrent cooling did ductivity water) on a laboratory scals are not increase the efficiency. J. Friedrichs and described by Bousfield (J.C.S. 1905, 87, 740), H. von Kroska (Chem. Fabr. 1934, 7, 281-287) mads tests on ten types of apparatns. The results are tabulated and discussed. The West apparatus (Ind. Eng. Chem. [Anal], 1930, 2, 199) gave best results. E. J. Williams (School Sci Rev. 1934, 16, 271) recommends double-aurface condensers for student and general laberatory use

For condensing vapours in high vacuum work a highly specialised type of "trap" condenser is used. The vapour trap generally consists of a U tube immersed in a refrigerant contained in a Dewar vessel (Fig. 15A) A



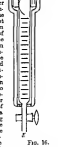
useful design, believed to have been origi-nated by S. Dushman, is shown in Fig. 15B, in which the vessel is unsilvered (see Hickman, J. Franklin Inst. 1932, 213, 119-154. The refrigerant (usually solid CO, or liquid air) be used, eccording to is placed in a, where it is insulated from the the temperature desurroundings by the operating vacuum, The vacuum line connects at b and e. This type of trap presents a large condensing surface and metal and contains a Imposes little resistance to gas flow.

The condensers so far described are usually constructed of glass, but occasionally metals and dividing the tube constructed of glass, but occasionally metals and dividing the tune are employed. The use of all adminishms for the pint for comparting the pint of pi numerous ammonia distillations, prefer a block- metal condenser tube is connected to the glass tin condenser. A convenient method of con-structing a number of such condensers in series structing a number of such condensers in series. This provides an air tight scal between the to be cooled by one stream of water is described metal and glass. The condenser is adapted to by Barnsrd and Bushop (J. Amer. Chem. Soc. 1906, 28, 999).

tested condensers by using them as reflux | enclosed in a metal tank which can be filled with apparatus; the loss of ether vapour under cold water; these condensers find considerable similar conditions was measured for each use in small plant for preparing distilled water. Golding (J.S.C.I. 1906, 25, 678), and by Hartley, Campbell, and Pools (J.C.S. 1908, 93, 428) The use of silica tubes for such condensers in now more common.

A condenser system made of glass and metal suitable for amalianalytical fractionating columns has been designed by Means and Newman (Ind.

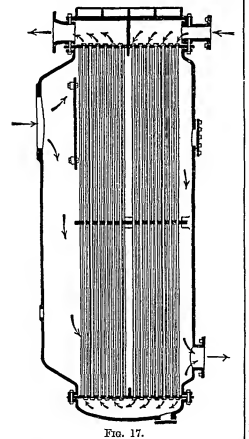
Eng. Chem [Anal.], 1936, 8, 231). It greatly resembles the top of a Prodbielniak column (sbid. 1931, 3, 177). An evacuated glass jacket, silvered efficient mediation, surrounds the burette and condenser. The condenser itself is a metal cylinder fitting into the top of the glass jacket and is insulated from the glass by means of an asbestos cord. The cooling medium is run down into the condenser through tube (Fig. 16), and escapes at D. Compressed carbon oxide or liquid eir can eired. The inner or condensing tube is of apiral strap extending down to the burette



burette by soldering to the platinised glass. distillations under reduced pressure, the vacuum connection being made at C and liquid extracted A nseful metallic condenser is the ball type, at E by any of the conventional methods for shown in Fig. 10B, used on the refux principle extraction of liquids during vacuum distillain numerous experiments involving extractions tions. Lattleton and Bates (Trans. Amer. Inst. with volatile solvents. Storeb's metallic redux Chem. Eng. 1926, 47, 95-100) have discussed condenser is depicted and described in J.S.C.I. the relative ments of Pyrex glass and metal condensers. The overall coefficient of heat The spiral (worm) type of condenser is fre- transfer from cooling water to vapour is not of quently constructed of metal (copper) and the same order of difference as would be expected

from the great difference in heat conductivity. The authors point out that an important factor in heat transfer is the film on either side of the cooling tube, the film itself acting as an insulator. This film is present on both Pyrex and metal condensers. (The effect of films is discussed below.)

In the design of large-scale industrial surface condensers stress is placed on efficiency. They are fundamentally heat exchangers designed for a special purpose. To secure a high coefficient of heat transfer from the condensing vapour to the cooling medium, Walker, Lewis, and McAdams ("Principles of Chemical

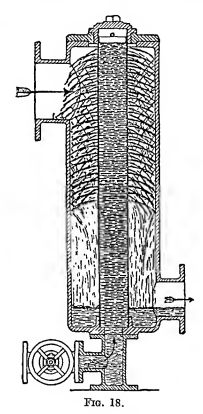


Engineering," 2nd ed., p. 398, McGraw-Hill, New York) state that the requirements are: "... rapid circulation of the cooling medium to reduce the effective thickness of the water film through which heat must flow by conduction, immediate removal of the condensed liquid, and as complete removal as possible of non-condensable gases to prevent the formation of an insulating film of relatively stationary Distillations done in vacuum help to meet the last requirement. A typical industrial surface condenser is shown in Fig. 17.

A surface condenser has the following advan-

vacuum distillations it is not necessary to pump the water out of the system or remove it by means of a barometric draw-off.

A parallel jet condenser is shown in Fig. 18 and a fall-pipe jet-condenser in Fig. 19. Jet condensers have a large capacity per unit floor space, they are of simple construction, of low cost, and corrosive vapours may be condensed without destruction to the chambers. However, a large volume of cooling water is needed, and in vacuum operations the water must be removed from the system either by pumps or through a barometric column. Gases dissolved in the water are also liberated directly



into the vacuum space and must be removed by the vacuum pumps.

Industrial condensers of various kinds are described in the following: Hausbrand, "Evaporating, Condensing, and Cooling Apparatus" porating, condensing, and cooling apparatus (trans. by Wright and rev. by Heastie), Benn, London; Sauvage, "Production et Condensation de la Vapeur," Baillière et Fils, Paris; Mironneau, "Technologie du Froid. T. I. Compresseurs et Condenseurs," Eyrolles, Paris; Low, "Condensers," McGraw-Hill, New York.

Other industrial condensers are described by the following: I.G. Earbenind A.G.

by the following: I.G. Farbenind. A.-G., Apparatus for Condensing Vapours by Streams of Liquid Passed in the Same Direction," B.P. 301430, 1927; Kalindustrie A.-G., "Surtages: the condensate is obtained unconface Condensers with Rotating Internal Cooling taminated with the cooling water, and in Drum," B.P. 306106, 1928; Leach, "Apparatus re issues 1437:1-1937s, 1935; Lach, Heat Exchange Apparatus Suttable for Condenside (Condenside Condenside Con "Heat Exchange Apparatus Sutahle (Resat Severe Corrosive Conditions," Condensing Oil Vapors," U.S.P. 1987051, Met. Eng. 1935, 42, 370-371.

1935; Cameron, "Condensing Hydrocarbon The foundation for the treatment as a Vapors such as those of Gasoline," U.S.P. escence of condensers and condensate. 1992352, 1935; McDermet (to Elhot Company), Spray Device Suitable for Use with Con-



Fig. 19.

densers, Evaporators, etc.," U.S.P. 2012454, 1935; Byer, "Counter eurrent Vapor Con

denser," U S P. 2013029, 1935. Some of the metals.

such as zme, lead, and cadmium, which are sufficiently volatile, are purified by evaporation and condensation Several of the apecial types of apparatus for condensing metal vapours have been designed by Fried, Krupp Grusonwerk A G., "Rotary Condenser for Metal Vapours," F.P 774132, 1935, Handwerk and Mahler (Assrs to The New Jersey Zine Com pany), "Apparatus and Method for Condensing," 421026, 1934; Fried, Krupp Gruson-werk A. G., Apparatus for the Fructional Conof Metal Vapours," BP. 427233. 1935 : G P. 608800, 1935. Many articles and

many patents have been issued on the prevention of corrosion in metal condensera. Special alloya have been developed for use in making condenser tubes sine

from the cooking water must be combatted as well as corrosion from the vapours. The preventive methods and the slloys have been discussed by: White, "Corrosion Tests on cuscussed by; White, "Corroson Tests on Condenser Tubes," Truns. Amer. Soc. Mech. Eng. (Fuels and Steam Power), 1931, 53, No. 14, 247-251; Worthungton, "Copper-Nickel Tubes. Their Advantage for Steam Condensers," Metal Progr. 1933, 24, 20-21; Meurk, "Protecting Surface Condenser Tubes from Corroson." U.S.P. 1978106, 1935; Korany and Bluss, "Prevention of Corroson in Gas Condensers," Mediagram, "Canada Condensers," Gas Aev Record, 1933, 78, 23, 24. densers." Gas Age Record, 1935, 75, 33-34; Tubes as a Function of their Brinell Hardness," Warme, 1935, 58, 173-177; Stager, "Topo- tions of heat-transfer coefficients on vertical tubes

for Oil Refining and Condensing," U.S.P. chemical Viewpoints" (corrosion tests on brass re issues 19377-19378, 1935; Leach, "Heat condenser tubes are described), Korrosion u

The foundation for the treatment as an exact ecience of condensers and condensation processes was laid by Molher (Z. Ver. deut. Ing 1897, Nos. 6 and 7). The first complete treatise was written by Hausbrand (1899), quoted above. Many articles and papers have followed since then. H. Braidy (Ind. Chim. 1919, 6, 201-205, 234-237) presented a mathematical discussion of the theory of condensers and of cooling devices in general. Brewer and Stivers (Mech. Eng. 1921, 43, 672-673) discussed the relative efficiencies of various types of cooling apparatus used as industrial units. Some features in condensing plant operation are given by Hardy (Proc. Univ. Durham. Phil. Soc. 1921, 6, 11, 66-83). The theory of fractional condensation was treated by Leslie (Ol News, 1924, 12, No. 3, 15–16) Lewis (Amer. Gas J. 1928, 128, No. 3, 42–44) has considered the cooling of non-condensable gases containing vapours which condense during the cooling process. Lewis discussed this case theoretically and showed that the base of condensation was and showed that the heat of condensation is not transmitted through the gas film on the pipes and should not be considered in calculating the coefficients. The vapour passes through the film by diffusion and is condensed on the cold pipe surface (see also E. F. M. van dar Held, Physics, 1934, 1, 1153-1160, "The Effect of Air on the Condensation of Water Vapour") Monrad and Badger (Ind. Eng. Chem. 1930, 22, 1102, 1112) 1103-1112) brought to light several important papers on heat transfer, especially that of Nusselt (Z. Ver. deut. Ing. 1916, 60, 541) When Nusselt's equations for heat transfer in the condensation of saturated vapour on the outside of horizontal tubes were applied to the work of McAdams and Frost (Ind. Eng. Chem 1922, 14, 13-18), Morris and Whitman (thid 1928, 20, 234-240), Clement and Garland (Univ. 111, Eng. Expt. Sta Bull No. 40, 1909), and Othmer (Ind. Eng. Chem. 1929, 21, 576-583), the experimental results agreed fairly well with the calculated figures When applied to data on vertical tubes (Badger, Monrad and Diamond, Amer. 1nst. Chem. Eng. (Detroit meeting), 1930, 23 pp, preprint), the experimental results were several times higher than the theoretical. The authors attempted to explain this by calculating the point on a vertical tube where turbulence hegins in the film of condensate. Nusselt assumed this film to be in viscous flow only. It was shown that turbulence hegins a relatively short distance down the tube, and this may account for the high results as compared with Nusselt's formula. A mathematical analysis similar to Nusselt's, but assuming turbulence in the film, was not made. Siegel, "The Corrosion Resistance of Condenser The variations in the degree of turbulence explain the great difficulty in making determinaA general review of heat-transfer processes Ind. Eng. Chem. 1921, 13, 819-820; Mach and which could be applied or are in actual operation Lederle, Chem.-Ztg. 1921, 45, 779. has heen given by Othmer (Ind. Eng. Chem. 1930, 22, 988-993). The article notes such important developments as mercury diphenyl heating, forced circulation evaporators, and high-pressure equipment (see also Badger, Monrad and Diamond, I.c.).

The effect of ribbed surfaces on the heattransfer rates in condenser sections has been discussed by Olson and Wentworth (Oil and Gas J. 1923, 31, Nos. 6, 14, and 16) and hy Du Pont (Natl. Petroleum News, 1923, 24, Nos. 46,

74-75, and 77-78).

Other papers on heat-transfer in condensers have been written by: Colburn and Hougen (Ind. Eng. Chem. 1934, 26, 1178-1182; Chem. Age, 1935, 32, 579-580); Heastie (Food, 1934, 4, 110-112; 1935, 5, 163-165); Frenc (Refiner and Natural Gasolinc Mfr. 1934, 13, 390); Pavloy and Kravetz (Khim. Mashinostroenie, 1935, No. 4, 19-21); Du Pont (Natl. Petroleum News, 1935, 27, No. 23, 110-111); Oldham (Ice and Cold Storage, 1934, 37, 27-28).

In the realms of physical science, the word condensation has assumed a broader meaning, which includes the condensation on charged particles in the Wilson cloud chamber, the formation of fog, the vesicular state, and the adsorption on various types of surfaces, particularly capillary adsorption phenomena. These special processes are discussed under headings of adsorption and theories of adsorp-

tion (v. Adsorption).

Appendix.—Other forms of the Liebig condenser have been described: Ber. 1892, 24, 3950; *ibid*. 1895, 28, 2388; 1899, 18, 707; Chem.-Ztg. 1902, 2 J.S.C.I. 26, 325; ibid. 1902, 26, 633; ibid. 1904, 28, 686; Bull. Soc. chim. 1904, [iii], 31, 1116; Chem.-Zig. 1904, 28, 598; *ibid.* 1905, 29, 809; B.P. 6916, 1905; Chem. Zentr. 1906, ii, 993; Editorial, L'industria Ind. Eng. Chem. [Anal.], 1929, 1, 43-44; Nanmann, Biochem. Z. 1930, 217, 409-420; Crandall, J. Lab. Clin. Med. 1930, 16, 89-91; Vaughn, J. Chem. Educ. 1931, 8, 2433-2434; Hanak, J. Prakt. Chem. 1933, 136, 202-204; Wagenschler, U.S.P. 1994934, 1935.

Other reflux condensers are described by: Chem. Ztg. 1896, 20, 462; Hahn, ibid. 1900, 24, 619; Bull. Soc. chim. 1901, [iii], 25, 476; Z. anal. Chem. 1901, 40, 769; Chem. Ztg. 1904, 28, 598; Chem. Zcntr. 1906, ii, 993; Chem. News, 1907, 95, 52; Bull. Soc. chim. 1908, [iv], 3, 855; Stoltzenberg, Z. angew. Chem. 1908, 21, 2084; Merkel, ibid. 976; J.S.C.I. 1908, 27, 21, 2084; Merkel, vind. 976; J.S.C.I. 1908, 21, 21, 2084; Merkel, vind. 976; J.S.C.I. 1908, 22, 362; Stolzenberg, Z. angew. Chem. 1909, 22, 351; Vollrath, Chem.-Ztg. 1910, 34, 1068; Haworth, Bertho and Siddiqui we owe the Anon., ibid. 716; Thorner, ibid. 1911, 35, 597; Heide, ibid. 531; Fishburn, J. Amer. Chem. Soc. 1917, 39, 1074; Vigreux, Bull. Soc. chim. Chem. Soc. 1932, 9, 553) the alkaloids in the 1917, [iii], 21, 46-48; Vigreux, Ann. chim. anal. chim. appl. 1919, 1, 211; Faust, Z. angew. difficult to extract with acidified EtOH or Chem. 1919, 32, 1, 183-184; Shohl and Koch.

Other double surface condensors are described by: J.S.C.I. 1905, 24, 1190; Chem. Zentr. 1908, ii, 277; Kob, Chem.-Ztg. 1910, 34, 116; Davies, ibid. 1911, 35, 531.

CONDURANGO. The dried hark of Marsdenia Cundurango Nichols (Gonolobus Cundurango Triana; Fam. Asclepiadaceæ). It is obtained from a half-climbing or prostrate shruh of Ecuador.

Kubler (Arch. Pharm. 1908, 246, 620) states that the principal constituents of the bark are dextrose, the glucoside condurangin, and a polyhydric alcohol conduritol. Condurangin, C40 H60O16, contains two methoxyl groups, is soluble in water, alcohol, and chloroform, insoluble in ether. Dilute sulphuric acid hydrolyses it to glueose and an amorphous powder, containing several substances; alcoholic potash yields some cinnamic acid. Conduritol, C6H10O4, erystallises from hot alcohol in colourless prisms, m.p. 142-143°, and yields a tetrahenzoyl derivative. According to Carrara (Gazzetta, 1891, 21, i, 204; 22, i, $2\bar{3}6$), commercial condurangin can he scparated into two parts, one insoluble and the other soluble in water. The former is a white powder, $C_{20}H_{32}O_6$, m.p. $60-61^\circ$; and the latter a yellow substance, $C_{18}H_{28}O_7$, m.p. 134° . Hager (Handhook, 1900) has further investigated these substances and named them a- and b-condurangin; a-condurangin gives a green colour with Fröhde's reagent (conc. sulphuric acid and molyhdic acid), b-condurangin does not. Carrara (l.c.) also isolated a yellow powder, $C_{30}H_{50}O_2$, which he named conduran-sterin, m.p. 52°.

CONDY'S FLUID. A solution of sodium permanganate, NaMnO₄, and sodium manganate, Na2MnO4

CONESSIDINE, CONESSIMINE CONESSINE AND MINOR ALKALOIDS FROM HOLARRHENA SPP.

CONESSINE AND MINOR ALKA-LOIDS FROM HOLARRHENA SPP. Holarrhena antidysenterica Wall (Fam. Apoeynaceae) or Kurchi-bark is well known in India as a remedy for dysentery as substitute for the more expensive ipecacuanha root. The plant contains about a dozen alkaloids which were partly isolated also from other Holarrhena spp., viz. H. africana D.C., H. congolensis S. Also Wrightia zeylanica contains similar alkaloids; it was from this plant that Haines first isolated conessine, also called wrightine in earlier papers (Trans. Med. Soc. Bomhay, 1858).

A survey of early literature is given in a paper by Kanga, Ayyar and Simonsen (J.C.S. 1926, 2123). Early investigators mostly isolated conessine, which is the chief constituent of the

Chem. 1919, 32, I, 183-184; Shohl and Koch, H2O. On the same hark all hitherto known

methods were tried and yields varying from 0.5 | given in detail they isolated three new alkaloids. to 1.7% crude total alkaloids were obtained.

EXTRACTION (Siddiqui et al., Ic.).-Dry powdered hark is percolated with a mixture of 80 parts of Et.O and IO parts EtOH, shaken with IO parts of NH,OH. The total alkaloids are precipitated by passing gaseous HCl through the percolate. The precipitate is dissolved in H.O and sodium sulphate is added when a first fraction of "insoluble sulphates" is obtained (0 17% calculated for free base). The motherliquor from the insoluble sulphates is made alkaline and exhausted with ether, when 21% "ether-soluble bases" are obtained A fraction of this ether residue is soluble in light petroleum and is subsequently precipitated from the moist petroleum with CO₂, while a fraction remains in the solvent. The bases insoluble in light petroleum are fractionated from CH, CO, Et with CO, Finally three main fractions are obtained

(a) Sulphates insoluble in H2O.

(b) Carbonates insoluble in light petroleum.

(c) Bases soluble in light petroleum, Fraction (c) (tertiary bases) gives 04-05% conessine; last traces of other alkaloids are removed by the "light petroleum carbonate method." Concessine may be purified by recrystallisation of its hydrogen oxalate (Pyman, J.C.S. 1919, 163; Kanga, Ayyar and Simonsen, fe.). From the mother liquors of the exalate, the hydrobromides are prepared and the base fractionally precipitated from this salt, the middle fraction yields isoconessimine.

Fraction (b) contains secondary hasea which are stronger than conessine. The bases were frectionally procepitated with ammonia, the first fraction containing "insoluble sulphates" and a fraction of "soluble sulphates." The third frection contains some more concesine and chiefly "carbonates insoluble in light petro-leum." The latter are united with the middle fraction and the hydriodides precipitated with KI (0.3%). After recrystallisation from MeOH and EtOH the carbonate is precipitated from light petroleum solution, the free base prepared and recrystallised from moist CH, CO, Et, when bars, easily soluble in McOH, less so in H, O or pure concessiman (0.02%) is obtained A later EiOH: B HgCl,+2HCl needles, sparneyl description (Siddiqui, J. Indias Chem. Soc., soluble also in hot H, O: B dimetholide, 1934, 11, 284) recommends dissolving fraction (b) in moist CH, CO, Et and precipitating the more atrongly basic di secondary consmine as carbonato whilst conessimine and isoconessimine remain in solution and are separated from each other by means of fractional crystallisation from CH, CO Et in which isoconessimina is more soluble than conessimme.

Fraction (a) is treated with 10% HCI, the white granular residue is dissolved in 10% alcoholic HCl and the sulphates precipitated from the hot neutralised solution with HISO4. The recovered bases are recrystallised from MeOH—CH₃-CO₂Et, when holarthine (0 02%) separates; from the concentrated mother-liquors holarthimine is obtained.

Bertho, von Schuckmann and Schönberger (Ber. 1933, 68, [B], 786) fractionated commercial mother liquors of the fraction soluble in light petroleum (Siddiqui's fraction c) with con easine as chief constituent). By a process not H.O.

conessidine, conkurchine, and kurchenine (yield of each alkaloid about 10% of conessine obtained). Haworth (J.C 2. 1932, 631), by extracting the seeds of H. antidysenterica, obtained conessing and norconessine. The two bases were separated

by the hydrogen oxalate method. Sudhamoy, Ghosh and N. N. Ghosh (J. Indian

Chem. Soc. 1928, 5, 477) extracted Kurch bark with EtOH. The total alkaloids (1.3%) were first precipitated with tartaric acid, when the tartrates of concessina and kurchicine separate. the two alkaloids are fractionated by recrystallisation of the sulphates and the free hases from CHCl, and light petroleum. The mother liquors from the precipitated tartrates contain kurchine.

Peacock and Chowdhury (J.C S. 1935, 734) exhausted Kurchi bark with EtOH; from the erude alkaloids (1%) they obtained an insoluble hydrochlonde fraction which yielded a new

alkaloid lettocine (0 1%). HOLARRHENINE is an alkaloid extracted with very dilute HC1 from the bark of H. congoleness (Pyman, i.e.); conessine (0.0%) was removed from the crude alkaloids by shaking out the alkaline solution with light petroleum. The alkaline mother liquors were in turn extracted with Et.O, when a small quantity of holarrhen-me was obtained.

The separation of the holarrhena alkaloids is extremely difficult and it is not clear yet whether the number of bases can be reduced owing to the fact that several of those isolated are identical (cf. Siddiqui, i.e. and Berthe et al , i.c.).

Conessine (I), C₈₄H₄₀N₈, [a]_D -19° (in CHCl₂) or +21-6° (in absointe EtOH), m.p. 126° crystallises from acctone in short needles or large prismatic plates, easily soluble in all organic solvents. (I) is a ditertiary base containing three NMe-groups; it is sensi-B 2HCI silky needles (from MeOH-acetone), [a] +9 32°, m p. 338-340°; B 2H1 colourless soluble also in hot H.O; B dimethiodide, slightly yellow octahedra, casily soluble in H.O or EtOH, m p. above 285°, it yields NMe, on distillation with AgOH.

Conèssimine (II) (Siddiqui et al., l.c.). C23H34Ng. [a] -22 25° (in CHCl3), mp 100°, microscopic needles (from dry light petroleum. CH. CO. Et oracetone), or m p. 91° (dihydrate), long slender needles (from moist CH, CO,Et). h p. 230°/I-8 mm. (II) contains two N—Megroups and one activa H-atom (Zerewitinoff). Mono - nitroso - compound, mono - henzoyl (11), m.p. 121°, monoacetyl (II) (oil). (11) yields (1) on methylation with HCHO and HCOOH; (1) therefore is mono-methyl (11). B carbonate unstable; B 2HCl amorphous, exceedingly soluble in H2O and EtOH, [a]20 -151', m p. 312°; B 2HI, m.p. 318-319° (from H₂O); B picrate, long rectangular plates (from EtOH) very sparingly soluble, in hot

Holarrhimine (III) (Siddiqui et al., I.c.), $C_{21}H_{36}ON_2$, $[a]_D^{25}$ -14·19°, m.p. 183°, crystallises from CH3·CO2Et in star-shaped radiating needles. (III) is very soluble in EtOH and CHCl3, almost insoluble in Et2O and light petroleum. Diacidic base, no OMe or NMe groups, three active H-atoms (Zerewitinoff). B·carbonate is unstable. B·2HCl, $[a]_D^{25}$ -22·80°, m.p. 345°, broad plates, easily soluble in EtOH, less so in H2O; B.2HBr thin plates and broad needles, m.p. 358-360°, sparingly soluble in H₂O; B picrate yellow plates, m.p. 198-200° (anhydr.), fairly soluble in hot H2O.

Holarrhine (IV) (Siddiqui, et al., l.c.), C₂₀H₃₈O₃N₂, [a]²⁵ -17·01°, m.p. 240°, crystallises from MeOH-CH3 CO2Et in needles. easily soluble in MeOH and EtOH, sparingly so in CHCl3, almost insoluble in CH3 CO2Et, Et₂O or light petroleum; (IV) appears to be a secondary base. B.picrate semi-crystalline

powder, m.p. above 320°.

isoConessimine (Siddiqui et al., l.c.) (V), $C_{23}H_{38}N_2$, $[a]_D^{28} + 30.0^\circ$ (in absolute EtOH), m.p. 92°, crystallises from light petroleum or acetone in needles, from moist CH₃·CO₂Et, as dihydrate, m.p. 88-92°; (V) is readily soluble in all organic solvents, it contains two NMegroups and one active H-atom (Zerewitinoff). (V) on methylation with HCHO and HCOOH yields (I), (I) therefore is monomethyl-(V); N-benzoyl-(V) m.p. 159-160° and N-acetyl-(V) m.p. 127-128° are monoacidic bases.

(V).earbonate is unstable, B.2HCl is erystalline, m.p. 335°, exceedingly soluble in H₂O or EtOH; B·2H needles, m.p. 316°, readily soluble in cold H₂O (difference from (I)); B-picrate, yellow rectangular plates and broad needles from dilute EtOH, m.p. 198-200° (anhydr.).

Conimine (Siddiqui, et al., l.c.) (VI), C22H36N2, $[a]_{\rm D}^{28}$ -30° (in absolute EtOH), m.p. 130°, crystallises from fairly concentrated solutions, Et,O, light petroleum CH₃ CO₂Et, or acetone in needles, from moist CH₃·CO₂Et as hydrate. (VI) contains two active H-atoms (Zerewitinoff) and one NMe-group. (VI) on methylation with HCHO and HCOOH yields (I); (I) is dimethyl-(VI), N, N-dibenzoyl-(VI) m.p. 250°, N, N-diacctyl-(VI) (m.p. 139-140°), N, N-dinitroso-(VI) m.p. 206-207°. B-2HCl needles, m.p. 318-320°, easily soluble in EtOH and H₂O; B·2HC intercets, soluble in EtOH and H₂O; B·2HI prismatic rods, m.p. 293°, fairly soluble in EtOH, less in H₂O; B·picrate, brilliant yellow prismatic rods from H₂O, m.p. 140-141°, very sparingly soluble in hot EtOH and H₂O.

Conessidine (Bertho *et al.*, *l.c.*) (VII), $C_{21}H_{32}N_2$, $[a]_D^{21}$ -52·2° (in CHCl₃), m.p. 123°, crystallises from acetone in thin needles or thick crystals. (VII) contains one NMegroup. B·2HI faintly yellow needles from H₂O, m.p. 259°; B·2HClO₄ thin needles from H₂O, m.p. 243° (decomp.).

Conkurchine (Bertho et al., l.c.) (VIII), $C_{20}H_{32}N_2$, $[a]_D^{21}$ -67.4° (in EtOH), m.p. 153°, small needles, easily soluble in CHCl3, MeOH, la hetcrocyclic ring-system.

EtOH, acetone, benzene, sparingly soluble in EtaO or light petroleum. (VIII) does not contain NMe-groups. Methiodide amorphous; sulphate m.p. 342°, oxalate m.p. 325°, and acetylcompound m.p. 233° are crystalline.

Kurchenine (Bertho et al., l.c.) (IX), $C_{21}H_{32}O_2N_2$, $[a]_D^{21}$ -92° (in 2N-HCl), m.p. 335-336°, crystallises from MeOH in thin leaflets, insoluble in Et₂O, sparingly soluble in MeOH and EtOH. (IX) does not contain OMe nor NMe-groups. B-sulphate crystalline, $\begin{array}{l} [a]_{\rm D} - 78 \cdot 3^{\rm o} \ ({\rm in} \ {\rm H}_2 {\rm O}). \\ {\rm Norconessine} \ ({\rm Haworth}, l.c.) \ ({\rm X}), \ {\rm C}_{23} {\rm H}_{38} {\rm N}_2, \end{array}$

[a]_D +6.7° (in absolute EtOH) is an oil b.p. 240/0.7 mm., very readily soluble in all organic solvents, it contains three NMe-groups. Boxalate nodules, m.p. 225-227° from EtOH or H2O; B.2HCI slender needles from EtOHacetone, m.p. 340° (decomp.); B'dimethiodide pale yellow prisms m.p. 310-312° (decomp.).

Kurchicine (Ghosh et al., l.c., and Ghosh and Bose, Arch. Pharm. 1932, 270, 100), (XI) C₂₀H₃₆ON₂, [a]_D -11·4° (in CHCl₃), m.p. 175°, crystallises from CHCl₃-light petroleum in needles, easily soluble in EtOH and CHCla, sparingly soluble in benzene and acetone, insoluble in light petroleum. B·2HCl ncedles, easily soluble in H₂O, less so in EtOH, m.p. above 260°; B·H₂SO₄,2H₂O sparingly soluble in cold H2O, leaflets, m.p. above 270°.

Kurchine (Ghosh et al., l.c.) (XII), C23H38N2, $[a]_{\rm D}^{32}$ -7.57° (in CHCl₃), m.p. 73-75°, b.p. 233°/1 mm., crystallises from Et₂O in needles, easily soluble in most organic solvents. B.2HCI, H₂O rhombic needles from H₂O, easily soluble in H₂O, less so in EtOH, decomp. about 220°; B.2HBr needles from H₂O, easily soluble in H₂O and EtOH; B·H₂SO₄ flat needles from H₂O, sparingly soluble in cold H₂O, decomp. at 270°.

Lettocine (Peacock and Chowdhury, l.c.) (XIII), C₁₇H₂₅O₂N, light-brown, microcrystal-line powder (from CHCl₃-light petroleum), m.p. 350-352°, soluble in EtOH and CHCl₃, sparingly soluble in Et2O and light petroleum; B.picrate, m.p. 198° (from absolute EtOH); B.methiodide, m.p. 235° (from hot MeOH). Does not yield an acetyl-compound.

Holarrhenine (Pyman, *l.c.*) (XIV), C₂₄H₃₈ON₂, [a]_D -7·51°, m.p. 197-198°, crystallises from CH₃·CO₂Et in silky needles and is readily soluble in EtOH and CHCl₃, l.c.) (A... n. D. 197–198° sparingly so in cold CH₃·CO₂Et, acetone, or Et₂O. B·2HBr flat needles (from H₂O), m.p. 265-268°. (XIV) contains three NMegroups and yields an acetyl-compound which is still diacidic.

Constitution of Holarrhena Alkaloids.

-It has already been mentioned that (I) is obtained by introduction of one Me-group into (II) or (V) and of two Me-groups into (VI) (Siddiqui, l.c. 1934, 288, 290). By demethylation of (I) with BrCN (von Braun's method) (V) and (VI) are obtained (Siddiqui et al., J. Indian Chem. Soc. 1934, 11, 787). They conclude from their experiments that in (I) one N atom is purely aliphatic forming the group -NMe,; crystallises from Et₂O or EtOH-H₂O in the second N-atom as >NCH₃ is a member of

dihydro-conessine m p. 105° on hydrogenation (Spath and Hromatka, Ber. 1930, 63, 126; Osada, Amer. Chem. Abstr. 1928, 22, 429); bromine is added (Ulrici, Arch. Pharm. 1918, 256, 76) and with H.SO, and KIO, (I) may be synthesis of vanillin. oxidised to diory conessine m p. 294-295° (Warnecke, Ber. 1886, 19, 60; Ulrici, Lc., 71; Giemsa and Halberkann, Arch. Pharm. 1918, 256, 201). The Hofmann and Emde degradation of (I) was studied by Spath and Hromatka (i c , with information concerning earlier results) The first step of the degradation of (I)-dimethohydroxide removes the aliphatic N atom, npens the heterocyclie ring, and yields apoconessine, m p 69° (see Kanga, Ayyar and Simonsen, ApoConessine is then degraded by Emde's method, when a nitrogen free compound, C , H 300 m p. 74-76° is obtained Ovidation expenments did not show any results; no benzenering seems to be present in (1). Spath and Hromatka conclude that (1) contains four reduced carbocyche rings and a hetero cyclie group with a side chain containing a double bond. The double bond may however he The double bond may however be in the hetero-ring No further details about the constitution of the Knrchi alkaloids are known,

Physiological Action of Kurchi Alka-LOIDS -Bark and seeds of Holarrhena untidysenterica are used in India against dysentery. Acton and Chopra (Indian Med. Gaz. 1929, 64, 207, quoted by Ghosh and Bose, le) state that the total alkaloids of Kurchi-hark have an action Like emetine, but they are less expensive and exert no blood depressor action, the iodo bismuth compound of the total alkaloids is given orally (cf. also Brown, Brit. Med. J. 1922, I, 993) The physiological action of the hark and its extencts in the treatment of intestinal troubles may also be due to the fact that the tannates, as which the alkaloids occur, would remain undissolved in the acid juices of the etomach and pass on to the seat of the disease in the intestines (Siddiqui, 1c) Pure conessine gives unex-pectedly poor results; evidently some other alkaloid present in "conessine" is responsible for the antidysentery action.

Conessine does not influence bird malaria (Goodson, Henry and Macfie, Biochem. J. 1930, 24, 874), but even in very dilute solution it acts on tubercle bacili (Mersoner and Hesse, Arch. exp. Path. Pharm. 1930, 147, 339).

For action of norconessme, cf. White (J. Pharm. Exp. Ther. 1933, 48, 79) and of holarrhenine, ef. Burn (J. Pharmacol. 1915, 6, 305, quoted by Pyman).

CONGLUTIN. A globulin protein con-

tained in the seeds of lupins. CONHYORINE one of the several poison. ous alkaloids found in hemlock. v. Contyn

ALKALOIDS CONICEINE α-,β-, γ-, δ-, ε, and φ- v. COVIUM ALKALOIDS

CONIFERIN, C10H21O22Aq, mp. 185°, [a]p -67°, the ß glucoside of the fir tree, yields clucuse and conferyl alcohol when hydrolysed by

emulsin. It has the formula : C.H.,O.O C.H. (OMe) CH: CH CH,OH

(f) contains one double-bond and yields On careful oxidation with chromic and glucovanillin is formed which may be further oxidised to gincovanillie acid or reduced to glucovanilly alcohol. All three glucosides are hydrolysed by emulsin. Coniferin at one time was used for the

It has been synthesised by Pauly and Fener. atem (Ber. 1927, 60, [B], 1031) via glucocomferyl aldehyde, which can be reduced by the rather unusual method of adding it to a solution of sugar undergoing brisk fermentation.

It was discovered in the cambial sap of Lanz decidua Mill, and occurs in the saps of the consfers in general; it is found also in beetroot

and asparagus,

It crystallises in colourless rosettes of pointed needles, aparingly soluble in cold water. The aqueous solution has a slightly hitter taste. When moistened with phenol and concentrated bydrochloric acid, an intense blue colour is formed; it is soluble in concentrated sulphune acid with a red coloration.

E. F. A

CONTINE v. CONIUM ALKALOIDS.
CONTININE v. CONESSINE AND MINOS

ALKALOIDS FROM HOLARBHENA SPP. CONIUM ALKALOIDS. Comino is one of several poisonous alkaloids found in hemlock, Consum maeulatum Linn. (Fam. Umbellifern) (Fr. Grande Ciguë; Ger. Giftschierling) is a tall biennial of the temperate regions of Europe, Asia and America and was the essential ingredient in the potion administered to con-demned criminals by the Greeks. Besides d-and I comines, hemlock contains N-methyl dand Leonine, y consceine, conhydrine, and psendoconhydrine. Power and Tutin (J. Amer. Chem. Soc 1905, 27, 1461) found a similar mixture of alkaloida in fool's pareley (Athus

Cynapsum Lnn).

EXTRACTION.—According to Chemnitius (J. pr. Chem 1928, [u], 118, 25) contine is prepared from finely ground seeds (0 4-0 5% total alkaloids) by mixing with wood wool, moistening with 15% aqueous NaOH, and extracting four times with warm Et.O in a copper vessel The concentrated extract is acidified with 50% acetic acid and Et. O totelly evaporated. After cooling, solidified fat is akimmed off, and the acid solutions are washed with Et2O, excess of 35% NaOH as added, the alkaloids extracted with ether and finally fractionated in a current of hydrogen. The fraction boiling up to 165° 15 converted inte comme salts; at 166-174° "common purum" of commerce distils and the distillate above 174° is fractionated for

perature is not raised above 185° conhydrine is left behind. According to von Brann (Ber. 1905, 38, 3108) if the fraction boiling up to 190° (containing no conhydrine) is benzoylated, the unchanged tertiary N-methylconsines can be removed from the ethereal solution by dilute acid. The ethereal solntinu is concentrated and poured into light etroleum when most of the benzoyl-8 amino butylpropylketone formed by interaction of benzoyl chloride with y-coniceine is precipitated. The solvent is then distilled off and the residue

consinum purum and conhydrine. If the tem-

distil at 200-210°/16 mm., and are thus separated from any residual derivative of γ-coniceine Benzoyl-δ-aminobutylpropylketone and the benzoyl-coniines can easily be converted into the parent alkaloids by hydrolysis.

Pseudoconhydrine usually accompanies conhydrine, from which it can be separated as EtOH-insoluble hydrochloride (Löffler, Ber. 1909, 42, 116). For separation of conline and pseudoconhydrine, see Späth, Kuffner and Ensfellner (Ber. 1933, 66, [B], 591).

d-Coniine (I), $C_8H_{17}N$, $[\alpha]_D^{19} + 15.7^\circ$, m.p. -2° , b.p. $166-167^\circ/760$ mm., is a colourless, strongly alkaline liquid having a peculiar penetrating odour and a burning taste. (I) is slightly soluble in cold H2O, less so in hot (turbidity on heating); it is steam-volatile and easily soluble in EtOH, readily soluble in Et2O, sparingly so in CHCl3. Exposed to the atmosphere (I) slowly darkens and resinifies. (I) dissolves in CS2 forming a complex thiocarbamate (Melzer, Arcb. Pharm. 1898, 236, 701). B·HCl, large rhombs (from H₂O), m.p. 220°; B·HBr needles, m.p. thom H₂O₁, m.p. 220°; B-HBI needles, m.p. 211°; B₂·H₂PtCl_g, H₂O orange-yellow crystals from H₂O, m.p. 175°; B-AuCl₄, m.p. 75°; B-picronolate, m.p. 195·5° (Warren and Weiss, J. Biol. Chem. 1907, 3, 327). For dissociation of (I) and its salts, see Kolthoff (Biochem. Z. 1925, 162, 303).

I-Conline, $[a]_D^{21}$ -15.3°, occurs only in small amounts; it closely resembles its d-isomeride. Ahrens (Ber. 1902, 35, 1330) reports slightly different melting-points of the salts, which statement is not confirmed by Löffler and Friedrich (Bcr. 1909, 42, 107). *l*-Coniine may be obtained from conhydrine through β-coniceine (Löffler and Friedrich, l.c.).

N-methyl-d-conline (II), C₈H₁₆N·CH₃, $[a]_{D}^{243}$ +81·3°, b.p. 173-174°/760 mm., is a colourless oily liquid resembling confine which may be obtained according to von Braun's method (Ber. 1905, 38, 3108). Of the two methylconiines the d-form predominates. (II) is best obtained by methylation of (I) with HCHO and HCOOH (Hesse and Eichel, Ber. 1917, 50, Methyl-iso-pelletierine hydrazone yields racemic methylconiinc when heated to 150-170° with Na-ethoxide (Hess and Eichel, Bcr. 1917, 50, 1396, cf. also Hess, Ber. 1919, 52, [B], 1622). B.HCl, needles, m.p. 188°; B2·H2PtCl6, m.p. 158°.

N-methyl-1-conline, [a]_D -81.9°, shows a similar behaviour to (II); for further details, see Ahrens (l.c.).

Conhydrine (III), $C_8H_{17}ON$, $[\alpha]_D +10^\circ$, m.p. 121°, b.p. 226°/760 mm. crystallises in colourless leaflets from Et2O. (III) is a strongly basic substance, soluble in EtOH and CHCI, moderately so in H2O and sparingly in Et2O. N. benzoyl-(III), m.p. 132°. Dissociation constants of (III) (see Kolthoff, l.c.).

The salts of (III) are crystalline, the aurichloride,

small rhombs or prisms melts at 133°.

Pseudoconhydrine (IV), $C_8H_{17}ON$, $[a]_D$ +1I·0° (in EtOH), m.p. 105°-106° from dry other in slender needles, or m.p. 80° from moist ether in plates, b.p. 236-236.5° (for further

fractionated when the benzoyl-d- and l-coniines | details and for separation from (III), see Löffler, Ber. 1909, 42, 960). N-benzoyl-(IV), m.p. 132-133° (Spath, Kuffner and Ensfellner, l.c.). (IV) is a strongly alkaline base yielding crystalline salts, B.HCl, m.p. 212-213°; B.HAuCl, m.p. 133-134°; B₂ H₂PtCl₆ slender goldenyellow needles, m.p. 185-186°.

γ-Coniceine (V), C₈H₁₅N, optically inactive, b.p. 171-172°/746 mm., is a conline-like oil, strongly alkaline and almost insoluble in H2O. It is separated from (I) by von Braun's method (Ber. 1905, 38, 3108). (V) on reduction yields rac. (I). The salts of (V) are crystalline, B.HCl, m.p. 143°, hygroscopic; B.HBr, m.p. 139°; B₂·H₂PtCl₆, m.p. 192°.

Other coniceines, α -, β - (=allylpiperidine), δ -, ϵ -, and ψ -coniceines are synthetic products and have been obtained in various ways from (I), bromo- or iodo-(I), conhydrine and pseudo-conhydrine (for references, see Loffler et al., Ber. 1904, 37, 1879; 1905, 38, 3326; 1909, 42, 94, 107, 122, 929, 948, 3420). δ- and ε-coniceines contain a bicyclic ring-system.

CONSTITUTION AND SYNTHESES .- (I) is 2-npropylpiperidine, since by zinc-dust distillation of the hydrochloride, congrine, a-n-propylpyrid-(I) was first synthesised and ine, is obtained. resolved by Ladenburg (Ber. 1886, 19, 430, 2578; 1889, 22, 1403; 1906, 39, 2486) and was the first synthesis of a natural alkaloid; the method has been modified several times (Ber. 1909, 42, 94; 1917, 50, 139). More recent syntheses are recorded by Lautenschlager and Onsager (ibid. 1918, 51, 602) and Koller (Monatsh. 1926, 47, 393). For synthesis of rac. (I), see also Bergmann and Rosenthal (J. pr. Chem. I932, [ii], 135, 277), rac. (I) is also obtained by ring-opening of indolizidine with BrCN (Ochiai and Tsuda, Ber. 1934, 67, 1011; cf. also Diels and Alder, Annalen, 1932, 498, 43). Synthesis of (II) by methylation of (I), see above (Passon, Ber. 1891, 24, 1678; Wolffenstein, ibid. 1894, 27, 2611; Hess and Eichel,

l.c. 1917). Conhydrine (III).—Willstätter (Ber. 1901, 34, 3166) obtained l-piperidine-2-carboxylic acid by oxydation of (III) with CrO3 and suggested that the hydroxyl-group in (III) must be located in the side-chain. The correct formula for (III) was first proposed by Engler and Baucr (ibid. 1894, 27, 1779) and has received confirmation by the work of Hess and Eichel (ibid. 1917, 50, 1386) and Spath and Adler (Monatsh. 1933, 63, 127).

III.

Pseudoconhydrine (IV) is a hydroxycomine, century in Cornish copper mines, it has sines since treatment with HI yields an iodo (I) which been found in quite small amounts at a few on reduction forms d (I). The location of the other localities. hydroxyl group in the piperidine ring was first advocated by Löffler (i.e. 1909, 116); Spath, Kuffner, and Ensfellner (i.e. 1933) demonstrated that (IV) is 2-n propyl-5-hydroxy piperidine.

By removing the elements of H₂O from (IV)

\$\psi\$ consceine is obtained (Loffler, \$\epsilon\$ 1909, 122).

\$\psi Coniceine (V) -Beside its natural occurrence (V) may be obtained by interaction of (III) and P.O. or fuming HC! at 200-220°. Loffler and T.chunke (Ber 1909, 42, 915) suggest (V) to be \$\times^a\$ unsaturated comme. (V) has been synthesised by Gabriel (ibid. 1909, 42, 4059) by hydrolysing & phthalimido butylpropylketone.

For colour reactions of conium alkaloids, see Dilling (Pharm J 1909, 29, 34, 70, 102 Biochem. J. 1909, 4, 286) For separation of micotine and (1), see llout (Arch Pharm 1893, 231, 376) For microchemical reactions of (1), eec Colo (Plulippino J. Sci 1923, 23, 97), Rosenthaler (Arch Pharm 1927, 265, 319) and Wagenaar (B 1929, 869) For analytual data, ere Leichard (Pharm Centr 48, 385) Krayer Arch exp. Path Pharm, 1931, 162, 342) detects traces of (I) by micro sublimation of the hydrochloride

PHYSIOLOGICAL ACTION OF COMICM ALEA-LOIDS (Henry, "Plant Alkaloids," 2nd ed , p 38). -All alkaloids contained in hemlock are poisonous. They produce paralysis of the motor nerve terminations and stimulation followed by depression of the central nervous aystem. Respiration is generally accelerated and deepened at first but eventually becomes slow and finally ceases while the heart is still strong and consciousness has just disappeared. By introduction of a double linking as in y coniceme the toxicity is greatly increased, whilst by the insertion of a hydroxyl group, as in the conhydrines, it is much reduced.

Schl CONKURCHINE r. COVESSINE AND MINOR ALKALOIDS PROM HOLARRIESA SPP.

CONNELLITE. A rare mineral consisting of basic chloro sulphate of copper with the approximate formula

2CuCl, CuSO, 19Cu(OH), H,O,

corresponding to buttgenbachite (qw) with

L. J. S. CONOUINAMINE v. CINCHONA ALKA

LOYDS (this vol., p. 160c) CONQUININE v. CINCHONA ALKALOIDS

(this vol. p 169b) CONSTANTAN. A nickel copper alloy. CONTRAMINE. Diethylammonium di-

ethyldsthiocarbamate (Brilish Drug Houses, London), BPC CONVALLAMARIN and CONVAL.

LARIN. Glycosides found in Convallaria, the dried inflorescence of Consallaria majalis L. (hily of the velley) v. CARDIAC GLYCOSIDES CONVALLATOXIGENIN E, CARDIAG

GLYCOSIDES. CONVALLATOXIN v. CARDIAC GLYCO

CONVICINE. A glycoside found in vetch seeds, it yields on hydrolysis alloxantin, ammonia and a hexose.

CONVOLVULIN, C, H, O,, m p 150°, is the glycosidic resin from the Tubera Jalapa, known as jalap resin, and like the roots of several species of the Convolvulaces family, has long been used as a purgative.

Convolvatin, on dissolving in alcohol and reprecipitating with ether (Votoček and Valentin, Coll. Trav. Chim. Tchécoslovaquie, 1929, 1, 477) is separated into an ether soluble gly coside containing the zero sugar epirhamnose and an ether insoluble compound, rhamno convolvuhe send, C₅₂H₉₃O₅₂,7Aq, which is hydrolysed to glucose (4 mol), rhamnose (2 mol), and convolvulinoho acid which Votoček and Prelog (Coll. Trav. Chim. Tchécoslovaquie, 1929, 1, 55) have proved to be 2 l1-dihydroxy-palmitio acid. They have suggested that in the glycoside the hydroxyl groups each carry 2 glucose molecules and I rhamnose molecule united to form a treacchande. The seme hydroxy palmitic acid is found elso in turpethin, which contains the sugars rhodeose and glucose.

E F. A. CONVOLVULUS PSEUOOCANTAB RICUS, ALKALOIDS OF .- C. pseudoentidoreus Echrenia (Fam., Convolvulacen) u ur digenous to Middle Asia; it was first investigated by Orechoff and Konowalowa (Arch. Pharm 1933. 271, 145), who isolated four alkaloids Ground aceds are morstened with 25% ammonia and then percolated with benzene. The benzene is extracted with HCl, which after addition of ammonia is extracted with CHCl3. On evaporation the CHCl, leaves the crude, semi-solid alkalords (0 406%) This residue is taken up in absolute EtOH; by addition of alcoholic HCl a crude mixture of the hydrochlorides of convolume (1), convolumne (II), and convol vidino (111) is obtained The oily mother-liquor contains convolveme (IV). (I) and (II) are separated by the different solubility of their hydrochlorides, that of (II) being easily soluble in absolute EtOH. By recrystallising crade (!) sulphate in place of nitrate. It forms velvet from light petroleum, a small insoluble fraction is his crusts of minute hexagonal prisms with a obtained from which (111) was isolated (Orichoff rich blue colour. First observed in the 18th and Konowalowa, Ber. 1934, 67, [18], 1163).

10 kg. of seeds yield 153.5 g. (I)·HCl, 10.7 g. (II) HČI, 3.8 g. (III) HCI, and a small amount of (IV) (Orechoff and Konowalowa, Ber. 1935,

Convolvine (I), C₁₆H₂₁O₄N, m.p. II5° (from light petroleum), optically inactive, crystallises in colourless needles, easily soluble in EtOH, CHCl₃, and acetone, sparingly soluble in hot H₂O, Et₂O, and light petroleum. (I) is a strong base, the salts are crystalline: B·HCl, m.p. 260-261° from EtOH in which it is sparingly soluble; B.oxalate, m.p. 265-266° from EtOH; B.HNO₃, m.p. 212-213° m.p. 26I-263° EtOH; B-picrate, from needles from EtOH; B-chloroplatinate, m.p. 240-241° insoluble; B-chloroaurate, m.p. 217° from hot H₂O; B methiodide, m.p. 230-231° from hot H₂O. On alkaline hydrolysis (I) yields a mixture from which Orechoff and Konowalowa (1935, l.c.) isolated veratric acid and nortropine, which shows that (I) is veratroyl-nortropeine and has the following struc-

By methylating (I) at the secondary N-atom, convolumine (II) was obtained.

Convolamine (II), C₁₇H₂₃O₄N, m.p. 114-115° from light petroleum, optically inactive, crystallises in stout prisms, easily soluble in EtOH, acetone, CHCl3, and benzene, sparingly soluble in Et2O and still less in light petroleum and hot H₂O. The salts are crystalline: B·HCl, m.p. 237-239° from EtOH; B·picrate, m.p. 263-264°, decomp. plates from hot EtOH or H₂O; B-chloroplatinate, m.p. 216-217° needles from hot dilute HCI; B chloroaurate, m.p. 201-202° (needles from hot dilute HCI; B methiodide, m.p. 257-259° colourless needles from H2O. On hydrolysis with caustic soda (II) yields veratric acid and tropine, it has therefore the structure (II) above. The base has been synthesised by interaction of veratroyl-chloride with tropine (Orechoff and Konowalowa, 1934, l.c.).

Convolvidine (III), C₃₃H₄₄O₈N₂ (?), m.p. 2-193° (from EtOH), optically active, 192-193° crystallises in small colourless needles, easily soluble in CHCl₃, sparingly soluble in EtOH and acetone, insoluble in H₂O, Et₂O, and light petrolcum. The salts are only sparingly soluble, B-picrate, m.p. 229°-231°, casily soluble in hot acetone. On alkaline hydrolysis (III) yields vcratric acid and a base not yet identified (Orechoff and Konowalowa, 1935, l.c.).

Convolvicine (IV), $C_{10}H_{16}N_2$, oily base, b.p. $100-125^{\circ}/5$ mm. or $250-260^{\circ}/760$ mm. Becomes yellow when exposed to the atmosphere. Easily soluble in H2O and the usual organic solvents. B-picrate, m.p. 200-202° needles from hot H2O. The two N-atoms apparently have a different basicity.

the plant contain (I) and (II) in about the same amount. During vegetation (II) seems to be slowly transformed into (I) (phytochemical demethylation).

Schl. COOK'S ALLOY. An alloy containing

43% zine and 57% antimony.
COOPERITE. Platinum sulphide, PtS, as tetragonal crystals, with the unit-cell dimensions a 4.91, c 6.10A. As minute grains it appears to be the main platinum-bearing component of the concentrates from the Bushveld norite, Transvaal. When first described by R. A. Cooper (J. Chem. Met. Soc. S. Africa, 1928, 28, 283; 1929, 4, 268, 275) it was thought to be PtS₂ and isomorphous with sperrylite later H. Schneiderhöhn (1929) described it as orthorhombic and isomorphous with marcasite [FeS₂]; while one crystal described by H. R. Adam (1931) as cooperite has since been proved to be laurite [RuS₂]. This is a case where chemical analyses have been made on mixtures, and crystallographic determinations have been made on single grains picked out from the mixture. By examining each grain separately by X-ray methods it has been possible to select sufficient homogeneous material for chemical analysis (F. A. Bannister, Min. Mag. 1932, 23, 189). The name cooperite was first used in 1869 for a variety of serpentine; it is also a trade-name for an alloy, Ni-Zr-W,

etc., used for cutting tools.

CO-ORDINATION COMPOUNDS .-When used in a chemical sense the term co-ordination is applied to a mode of atomic linking first recognised in 1893 by Alfred Werner, who subsequently devoted the remaining years of his life to a systematic study of co-ordination com-pounds. The hypotheses proposed by him in explanation of the results of his researches are embodied in the co-ordination theory, which affords a simple and comprehensive explanation of the chemical constitutions of the most diverse types of complex metallic salts, including aquoand ammino- derivatives, mordant dyes, and many representative mincrals. Even before Werner's time numerous chemical compounds were known, the formation of which could not be explained on the older theories of valency. Such substances, often termed "molccular compounds," were generally formed from simpler compounds in which the combining elements had exercised their valencies in accordance with their position in the periodic classification. For instance, hydrogen chloride, a combination of univalent hydrogen and chlorine in atomic proportions, combines with ammonia, a compound of tervalent nitrogen with three univalent hydrogen atoms to form the molecular compound ammonium chloride. This secondary combination of hydrogen chloride and ammonia was formerly regarded as being due to the possession by these substances of a "residual affinity," a force which was then considered to differ from that of valency.

Werner examined a large number of metallic ammines, compounds of similar type to am-Orechoff points out that in the seeds 90% of monium chloride, although generally more the alkaloids present are (I), whereas the leaves of complex in molecular structure, and showed

that they could all be formulated on a new] principle. The underlying idea of Werner's diamine and radicals such as the oxidate group theory was that the combining powers of an C₂O₄, which each function in the co-ordination atom depended not only on the nature but also on the number of atoms or groups with which it could become associated. It is this number, which Werner called the "co ordination num ber," rather than the valency in terms of lydrogen equivalents, which gives a measure of the total combining power of the element. A general survey of complex metalise salts shows that the co ordination number is most frequently 6, sometimes 4, and less often 8 or other values.

In writing graphic formule in accordance with the co-ordination principle the convention is usually adopted of enclosing the central nuclear element and its associating units within a square bracket, which thus contains all the elements implicated in the co-ordination compley or inner sphere It should, however, be explained that this co-ordination principle supplements but does not supersede the obler conceptions of valency Any accurately written co ordination formula can be dissected into constituent parts which are expressible in terms of integral valencies The verification is readily seen in the case of ammonium chloride and is equally applicable in the following instances [Cr6NH₃]Cl₃ and K₄[Fe(CN)₆], where the generators of the complex ammine are CrCl₂ and NH, and those of the complex cyanido are KCN and Fe(CN). These two compounds furnish examples of the 3 types of stomic linkings recognised by chemists (i) a non polar, non ionised link as exemplified in the combination of nitrogen and hydrogen in am monia or in the attachment of carbon to nitrogen in the eyenide radical, (ii) a polar or ionised link between the oppositely charged ions of a salt as in dissolved CrCl₂ or KCN, and also between the complex cation [Cr6NH₂] and the 3 chloring ions or between the complex amon [Fe(CN)_k] and the 4 potassium ions, (iii) the co ordination link postulated by Werner as existing between chromium and 6 ammonia or between iron and 4 cyanogen groups.

In 1837 a remarkable foreshadowing of Werner's views was published by Gibbs and Genth in a research on ammomo cobalt bases wherein they referred to the cobalt atom as "hexatomic," although they also recognised that the combining capacity (principal valency) of this atom was 3 They referred to the inner co-ordination sphere of cobalt with 6 associating units as the "conjunct" ("Smith-sonian Contribution to Knowledge," Washing ton, 1857, IX)

MANIPOLD ASSOCIATING GROUPS.-In the examples, [CreNH,]Cl, foregoing Ka[Fe(CN)a], the ammonia and cianogen groups occupy only one position within the co ordination sphere, and a similar degree of association is manifested by the halogens, water. hydroxyl groups, oxygen, pyridine, methylamine, dialkyl- or diaryl sulphides, phosphines, arsines, and many other molecular species. But coordination is by no means confined to such single associating units. There are a great many examples of compounds which contribute 2, 3, or 4 units to the co ordination complex.

Chelate Groups .- Compounds such as ethylene sphere as two associating units or doubly bound groups, often to form exceedingly stable co ordination complexes, have been grouped by Morgan (J.C S 1920, 117, 1457) under the convement general term of chelate groups [from xnk]
(Greek), chela (Latin), chely, (English)] the crab's pincer claw Certain of these chelate groups tion of atereochemical structure of co ordination compounds.

2 2'-Dipyridyl (I) and o phenanthroline (II) furnish examples of such chelate groups, both of which were employed by Werner in the demon stration of octahedral structure.

Blau had previously prepared the dark red to 2 2' dipyridyl ferrous bromido

(Monatsh 1898, 19, 647), and Werner (Ber. 1912 45, 433) showed that this salt could be resolved muto optically active forms Morgan and Burstall have since resolved the analogous pals red nickel salts, [Ni3dipy]Cl, 6H, O, into dextro- and lavo-components (JC & 193), 2214). The corresponding orange red bivalent rathemum salt [Ru3dipy]Cla6HaO has sleo proved sufficiently stable for resolution into optically active forms (Burstall, J C S 1930, 173)

These 2 chelate bases have also proved unful in demonstrating the brialency of biler. Hicker and Muhlbaner (Ber. 1928, 61, [B]. 2149) base obtained byvalent silver salts such

[Ag2phenan]S.O. or X.,

where X=HSO4, NO5, ClO3, or ClO4, and Morgan and Burstall (JCS, 1930, 2594) have prepared the salts [Ag2d:py]S2O2 or (HSO); while Barbieri (Atti R. Accad, Lince, 1931. 16, 44) has obtained the derivatives

$[Ag2d_1py](NO_3)_2$ or $(ClO_4)_2$

Chelate groups need not be organic, for Werner was able to show that the basic cobaltic salt, dodecamminohevoltetracobaltie chloride containing 3 chelate groups {(HO)₂Co4NH₃} is resolvable into optically active forms (Ber. 1907, 40, 2113, 1914, 47, 3087).

$$\left[C_0\left(\frac{HO}{HO}\right)C_0\left(NH_3\right)_4\right]_3$$
CI

In more recent times, Mann (J CS. 1933, 412) has replated optically active components of a complex rhodium salt containing sulphamide

$$\left[\left\{SO_{2} \stackrel{NH}{\sim} \right\}_{NH} \stackrel{V}{\sim} \left(OH_{2}\right)_{NA} \right]$$
 Na

metal with formation of a stable five-membered | shown in the following series : ring system, the foregoing sulphamide derivative contains a four-membered ring. Other examples of this type of chelation are present in complex sulphates and carbonates such as

and
$$\begin{bmatrix} O_2 S & O & C_0 4NH_3 \end{bmatrix} CI$$

$$\begin{bmatrix} O:C & O & C_0 4NH_3 \end{bmatrix} CI$$

and also among many polynuclear compounds (v. infra).

Six-membered chelate rings are very plentiful among co-ordination compounds and include derivatives of 1:3-diaminopropane,

of malonic acid, HOOC·CH2·COOH, and the characteristic and notable metallic complexes of

acetylacetone.

Metallic Acetylacetones.—The univalent radical $CH_3 \cdot CO \cdot CH : C(CH_3) \cdot O$ —derived from a cetylacetone furnishes an important chelate group, and similar groups are derived from the homologous β -diketones and analogous β -keto esters. This acctylacetone radical enters into combination with many of the metals as a doubly bound group and gives rise to cyclic co-ordination compounds of remarkable stability, often having other exceptional properties such as unusual colour, solubility in organic media, and volatility without decomposition. In the last respect this cliciate group may be said to give wings to metals. Aluminium and scandium acetylacetoncs represented by the general formula (I) (where X'''=Al or Sc) are volatile without decomposition.

Thorium acetylacetone containing four of these chelate groups has also been distilled. Compounds of similar type have been obtained from zirconium, hafnium, and from quadrivalent cerium (II, where Y^{IV}=Th, Zr, Hf, or Ce^{IV}). The acetylacetone radical "Ac" and its homologues co-ordinate with the elements boron, silicon, germanium, titanium, tin and zirconium conferring ionic properties on the complex so that polar compounds arise. The boronium and siliconium compounds studied by Dilthey (Annalen, 1905, 344, 300; Ber. 1903, 36, 923, 1395, 3207), and Rosenheim, Löwenstamm and Singer (ibid. 1833) are of the types.

Whereas many chelate groups are attached to a other elements of the fourth periodic series are

$$\begin{array}{lll} & [\text{Ti''Ac''}_3]\text{FeCl}_4\;; & [\text{Ge''Ac''}_3]\text{CuX}_2\;; \\ & [\text{Ge''Ac''}_2\text{X}_2]\;; & [\text{Sn''Ac''}_2\text{X}_2]\;; & [\text{Zr''Ac''}_3]\text{Cl}\;; \\ & [\text{Zr''Ac''}_4]\;; & [\text{Hf''Ac''}_4]\;; & [\text{Ce''Ac''}_3]\;; \\ & [\text{Ce''Ac''}_4]\;; & [\text{Pb''Ac''}_2]\;; & [\text{Th''Ac''}_4]\;. \end{array}$$

Although the vast majority of chelate rings are those of five and six members, there are a few examples of seven- and even eightmembered rings. The former including carboxylic acids such as succinic, maleic, and itaconic acids, are exemplified by the cobaltammine (I):

$$\begin{bmatrix} \mathsf{CH}_2 \cdot \mathsf{CO} \cdot \mathsf{O} & \mathsf{Co}(\mathsf{en})_2 \\ \mathsf{CH}_2 \cdot \mathsf{CO} \cdot \mathsf{O} & \mathsf{I}. \\ \end{bmatrix} \mathsf{NO}_3$$

$$\mathsf{I}.$$

$$\begin{bmatrix} \mathsf{O}_2 \mathsf{S} & \mathsf{CH}_2 \cdot \mathsf{CO} \cdot \mathsf{O} \\ \mathsf{CH}_2 \cdot \mathsf{CO} \cdot \mathsf{O} & \mathsf{Co}(\mathsf{en})_2 \end{bmatrix}$$

$$\mathsf{II.}$$

$$(\text{"en"}=\mathsf{ethylenediamine.})$$

(Duff, J.C.S. 1921, 119, 385) whereas the eightmembered ring is present in the complex salt of sulphonyldiacetic acid (II) (Price and Brazier, J.C.S. 1915, 107, 1367).

Tridentate Groups .- Threefold associating groups have been recognised such as $\alpha\beta\gamma$ -triaminopropane="tpn," which co-ordinates with cobalt and rhodium as a tridentate radical to furnish the complex salts [Co2tpn]Cl3 and [Rh2tpn]i, (Mann and Pope, Proc. Roy. Soc. 1925, (A), 107, 80), and 2:2':2"- tripyridyl or di-2-pyridyl-2:6-pyridine ("tripy") which forms two series of salts containing respectively 2 and 1 molecules of the triamine. The former series is exemplified by the purple ferrous salt [Fe2tripy]Br₂ and 4H₂O and the latter by the stable red platinous compound

(Morgan and Burstall, J.C.S. 1932, 22; 1934, 1498).

Quadridentate Groups.—A fourfold associating unit is furnished by the bivalent radical of ethylcnediaminebisacetylacetone,

$$\{CH_2\cdot N: C(CH_3)\cdot CH: C(OH)\cdot CH_3\}_2=ec,$$

which forms remarkably stable complexes R¹¹ec with copper, nickel, and palladium (III).

In the absence of air the two former compounds may be boiled without decomposition. The cobaltic complex salts [Coec2NH₃]Br exist in stereoisomeric forms (Morgan and Main Smith, J.C.S. 1925, 127, 2030; 1926, 912). Tri-β-aminoethylamine N(CH₂·CH₂·NH₂)₃= and Singer (thia. 1833) are of the types $[B''Ac''_2]FeCI_4$ or $AuCI_4$; $[B''Ac''_2]PtCI_6$; $[Tri.\beta.aminoethylaminc] N(<math>CH_2\cdot CH_2\cdot NH_2)_3 = [Si''Ac''_3]FeCI_4$ or $AuCI_4$; $[Si''Ac''_3]_2PtCI_6$. "tren" also behaves as a fourfold associating Following on silicon the various modes of interaction with acetylacetone exhibited by $[PtCI_2]_2$ and $[PtCI_2]_2$ (Mann and

to the cobaltic derivative

[(SCN),Cotren]SCN,H,O

(Mann, J.C.S. 1929, 409).

Fourfold co-ordinating units of a different type are represented in the naturally occurring pigments chlorophyll and hæmin (v. snfra), which possess a constitution based on that of porphin. the parent of the series, which yields metallic derivatives of the type IV (Fischer and Gleim, Annalen, 1935, 521, 157; Rothemund, J. Amer Chem. Soc. 1936, 58, 625).

STEREOCHEMICAL DEMONSTRATION OF CO. ORDINATION.—The prevalence of co-ordination numbers 4 and 6 suggested that co-ordination compounds had a spatial errangement of their associating units corresponding respectively with the tetrahedron and octahedron. Moreover, many compounds with a co-ordination number of 4 possess a planar distribution of addends with the metallic atom at the centre

of a flet four-point figure
Octahedral Structure -- Werner predicted that cobaltemmines containing co-ordination complexes of the type [CoA,B,] would exist in two stereo isomeric forms, and that if the 4NHs groups of example [CoBr,4NHa]Br were replaced by 2 mols of ethylenediamine, "en" replaced by 2 moss of echylenculaturis, en-=NH₂'CH₂ CH₃ NH₃, then one of the two isomeno salts [CoBr₂2en]Br would be resolvable into optically active forms (enantiomers). His verification of the latter pre-diction in 1911 gained world-wide acceptance of the co-ordination hypothesis and raised it to the level of a general theory of chemical combination and molecular structure. This demonstration was first accomplished

chloroamminodiethylenediammocobaltic chloride [NH,CoCl2en]Cl, which exists in 2 isomeric modifications depending on the spatial arrangements of associating units in the coordination sphere as illustrated by the following figure :

Cas or 1:2 Form. Resolvable into enantiomerides.

The racemoid cis-chloride was converted into d-bromocamphorsulphonate when the dd-salt was found to be more sparingly soluble than the

Pope, Proc. Roy. Soc. 1925, 109, A, 455), and M-salts. The two d-bromocamphorsulphonates were then aeparately converted into bromides which in 1% equeous solution gave [a]c+43 corresponding with molecular rotation

(C line 656 3mu). These two optically active salts were very stable, showing no tendency to racemise (Werner, Ber. 1911, 44, 1887). Other interesting complex salts, which exhibit

optical activity, possess the structures [M3drpy]Cl,6H,O

(where M=Fe, Ru, Os, Ni, and dipy=2 2'. dipyndyl), [M3en]Cl, (where M=Co, Cr, Rh, and en=ethylenediamine) and

 $[M(C_*O_*)_*]K_*$

(where M≈Al, Cr, Fe^{III}, Co^{III}, Ir, and Rh). To these salts the chelate groups 2 2'-dipyndyl, ethylenediamine or oxalate radical span the six points of en octohedron with the metal st the centre.

Representative co-ordination compounds employed in subsequent demonstrations were the following: [Co2NH,2en]Br, [Co3en]Cl,; NH.

obtained in meso and racemoid [CrCla2en]X ; [Cr3en]X, ; [Cr(C.O.)] K1; [Rh3en]Cl. Optically active forms of compounds having octahedral symmetry have been isolated in the case of the following elements as the central atom : sluminium, arsenic, chromium, cobalt, iron, nickel, indium, platinum, rhodium, ruthenium and osmium.

In recent years, X-ray analysis has been increasingly used in the determination of structure among co ordination compounds and a considerable number have been shown to possess an octahedral distribution of valencies by this The compounds K,[Fe(CN),] means K,[PtCl,], [CoSNH,]Cl, K,[OsCI,NH,] [RhCl5NH,]Cl, represent octahedral structures determined by X-rays. As final examples of octahedral co ordination compounds the carbonyls of chromium, molybdenum and tungsten M(CO), (M=Cr, Mo, or W) may be cited since these compounds owe their existence entirely to co-ordinate links between carbon monoxide and the metal atom. The arrangement of addenda is indicated in the following figuro (cf. Hieber, Romberg and Mühlbauer, Z. anorg. Chem. 1935, 221, 321-349).

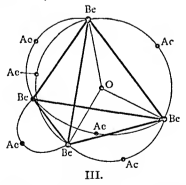
Tetrahedral Structure.—Even hefore the recognition hy Werner of the co-ordination link a demonstration of tetrahedral structure had been made in the case of elements other than carhon. In 1899 Pope and Peachey showed that quaternary ammonium salts of the type [Nabcd] + X could he resolved into optical antipodes. Amine oxides [NabcO] were obtained in optically active forms by Meisenheimer in 1908, and in 1911 this observer and Lichtenstadt prepared optically active phosphine oxides PabcO.

Since the enunciation of the co-ordination theory, tetrahedral symmetry has heen demonstrated in the case of several other elements. Borosalicylic acid (I) has yielded a strychnine salt with an optical activity due to the horon complex (Böeseken and Meulenhoff, Proc. K. Akad. Wetensch. Amsterdam, 1924, 27, 174); and Mills and Gotts resolved the heryllium derivative of benzoylpyruvie acid (II) and employed similar means in demonstrating tetrahedral symmetry in zinc compounds (J.C.S. 1926, 3121).

The tetrahedral distribution of associating units round oxygen is indicated by the work of Bragg and Morgan (Proc. Roy. Soc. 1923, 104, A, 437) on basic beryllium acetate,

$$Be_4O(CH_3 \cdot CO_2)_6$$
 (III)

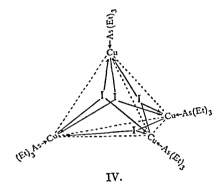
and its homologues (Morgan and Astbury, ibid. 1926, 112, 441).



Significant results have heen obtained from determinations of structure by means of X-ray analysis. A tetrahedral arrangement of molecules in cuprous compounds with a co-ordination number of 4 has been demonstrated for the well-known cyanide $K_3[Cu(CN)_4]$ and the thio-acetamide salt

$$\left[Cu \left(\leftarrow S = C \left(\begin{matrix} CH_3 \\ NH_2 \end{matrix} \right)_4 \right] CI$$

hy Cox, Wardlaw and Wehster (J.C.S. 1936 775) whereas 4-covalent cupric derivatives (v. infra) are planar. A structure of novel type has heen found hy Mann, Purdic and Wells (J.C.S. 1936, 1503) for the compound of cuprous iodide with triethylarsine, [(Et)₃As -> CuI]₄. The arrangement is tetrahedral with copper atoms at the four vertices of the figure whereas the iodine atoms are centred hut situated rather ahove the four faces of the tetrahedron (IV).

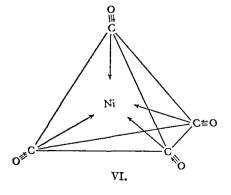


Univalent silver derivatives with a co-ordination number of 4 are also tetrahedral as exemplified by

$$\left[Ag\left(\leftarrow S=C \begin{pmatrix} CH_3 \\ NH_2 \end{pmatrix}_4\right]CI$$

Whilst Hein and Regler (Ber. 1936, 69, [B], 1692) obtained indications of optical resolution with the silver derivative (V) of 8-hydroxyquinoline,

which must, therefore, be tetrahedral. Other 4-covalent compounds of tetrahedral structure are the cyanides $K_2[M(CN)_4]$ (M=Zn, Cd or Hg) and the well-known and technically important nickel carbonyl Ni(CO)₄(VI) (Sutton, New and Bentley, J.C.S. 1933, 652; Brockway and Cross, J. Chem. Phys. 1935, 3, 828).



Planar Structures .- In the foregoing survey of molecular structure among co ordination compounds it was noted that when manufesting their fundamental univalency, copper and silver exhibited a tetrahedral arrangement of addenda in their fourfold co-ordinated compounds. It has now been found that in their higher valent states, copper, silver, and gold exhibit a planar configuration when the co-ordination number is By X-ray analysis cupric acetylacetone, Cu(C₅H₁O₂)₂ has been shown to be planar (Cox and Webster, JCS 1935, 731) and the com pounds [CuCl,,2H,O] and [CuCl,2py] have also been shown to be similarly constituted (Cox, Sharratt, Wardlaw and Webster, J.CS 1936, 129; Harker, Z. Krist. 1936, 93, 136). The planar arrangement among bivalent silver derivatives has been demonstrated by X ray atudy of the argentic derivative of picolinic acid (Cox. Wardlaw and Webster, J C.S 1936, 775)

and torvalent gold in potassium auribromide K[AuBr4]2H2O has a planar distribution of d bromine atoms round it (Cox and Webster, J.C.S. 1936, 1633)

Among tin and lead compounds, it is the lower valent compounds such as K, [SnC!,]2H,O and [PbCl,2SC (NH,),] which have a planar structure in contrast to the tetrahedral arrangements of stannic derivatives represented by SnI, and Pb(C2H2)4 in which a fundamental quadri Valency is manifested (Cox, Shorter and Wardlaw, Nature, 1937, 139, 72) A con-alderable amount of work has also been earned out on planar compounds containing elements of group VIII in the Periodic Classification. Chemical evidence in support of planar con figurations has been obtained by the isolation of ers and trans- forms of the nickel and palladium talta vi. bernylmethylglyoxane (i. and M) (Sugden, J.C.S. 1932, 246; Dwyer and Mellor, J. Amer. Chem. Soc. 1935, 57, 605).

The complex dithio oxalates of nickel, palladium and platinum (III) have been shown to be planar
by X-ray analysis (Cox, Wardlaw and Webster,
LCS. 1935, 1475). Co-ordinated birdent
by the discovery of the phthalocyanines, a erict

platenum derivatives have, in some cases, been Platform derivatives are the inception of the Coordination Theory by Alfred Werner who concluded that the two platesammines (PtCl,2NH,) were cis- and trans. isomerides

Recent X ray analyses support this conclusion and extend planar configurations to K.IPtCl.1. [Pt4NH₃]Cl₂H₂O, [PtCl₂,2SMe₂], and many other compounds including the corresponding palladium derivatives. Chelated platinammines also conform to a planar distribution of valences, since Drew and Head (J.CS 1934, 221) here obtained the two isomeric dichlorides :

A similar pair have also been prepared contain ing 2 mols. of isobutylenediamine

Another elegant proof of the planar arrangement of addenda emong platinammines has been furnished by Mills and Quibell (J.C S 1935, 839), who prepared a salt with the following cation:

and have resolved it into optically active forms which could not exist if the ion were tetrshedral. A final example of planar platinous saits is a derivative containing 2.6 di 2' pyridyl pyridine. in which a tetrahedral arrangement of the tridentato addendum is seen from the following figure to be madmissible (Morgan and Rurstall, J.C.S 1934, 1783).

stability, conveniently produced hy

M=Cu, Be. Mg, Ca, Zn, Cd, Ba, Sn^{II}, Pb, Fe, Mn, Co, Ni, Pt, AlCl, Vo, Sn^{IV}Cl₂.

heating the metal or its chloride with phthalonitrile (1:2-dicyanobenzene), they suhlime, in many cases, without decomposition at high temperatures. The planar configuration of the phthalocyanines is confirmed by X-ray examination. The foregoing formula suggests that the organic molecule functions as a fourfold associating unit (quadridentate group). The copper derivative is an important pigment now known commercially as Monastral Fast Blue B.S. (Linstead and co-workers, J.C.S. 1934, 1016–1033; 1936, 1719–1738). The corresponding naphthalocyanines have also heen prepared from 1:2-dicyanonaphthalene (idem J.C.S. 1936, 1744).

POLYNUCLEAR COMPOUNDS. — Co-ordination compounds may contain 2 or more nuclear atoms in the complexes. Among non-polar substances are hasic heryllium acetate Be4OAc6 and its homologues. From a very large group of complex polynuclear salts the following comparatively simple examples are selected:

$$\begin{bmatrix} en_2Cr(OH)_2Cr \ en_2 \end{bmatrix}Br_4; \\ \begin{bmatrix} NH_2 \\ OH \end{bmatrix} Co(NH_3)_4 \end{bmatrix} (NO_3)_4; \\ [Cu_25tu](NO_3)_22Aq.; [Cu_37tu](NO_3)_3; \\ [Cu_25etuH_2O](NO_3)_2,3Aq.; and \\ [Ag_23etu](NO_3)_2; \end{bmatrix}$$

where "tu"=thiourea and "etu"=ethylene thiourea. A series of remarkable chloroacids has the general formula [H₂O·X₃Cl₇]H,3Aq. where X=Mo, Ta, or W (and prohably Nb). These complex acids are derived from the chlorides (XCl₂)₃ or X₃Cl₆, which contain a stable inner core X₃Cl₄. Accordingly the structure structure

$$CI\left\{-X \underset{CI}{\swarrow} X \underset{CI}{\swarrow} X - \right\} CI$$

has been proposed for the parent chlorides. The complex phospho-molyhdic and tungstic acids are limiting examples of a numerous and diversi- TION.—The electronic conception of matter

of metallic complexes of unusual structure and | fied class of hetero-polynuclear acids ("heteropolyhasic acids"); their salts may he written as [PO₄12XO₃]R₃, nAq., where X=Mo or W and R=Na, K or NH₄. Moreover, phosphorus may he replaced hy arsenic. These derivatives contain a hasic tetrahedron with phosphorus at the centre and 4 oxygen atoms at the points of the solid figure. On this simple structure is huilt up a symmetrical arrangement of 12 interlocking WO a units which are divided up into 4 groups of 3 octahedra. In each WO₆ group, I oxygen atom is held in common with the phosphorus atom (and is also shared with 2 other WO₆ groups), 2 are shared with tungsten atoms of its own group of 3 octahedra, 2 with metallic atoms of other groups of octahedra, and 1 is unshared by other atoms. A arrangement is exhibited hy the acids $H_4[MX_{12}O_{40}]$, $H_5[BW_{12}O_{40}]$ and $H_6[H_2W_{12}O_{40}]$, where M=Si, Ge, and X=Mo and W (Keggin, Proc. Roy. Soc. 1934, 144, 75; Illingworth and Keggin, J.C.S. 1935,

> OTHER CO-ORDINATION NUMBERS.—In addition to co-ordination numbers 4 and 6, which suggest respectively tetrahedral and octahedral arrangements, there are metallic complexes and complex salts containing 8 associating units such as the acetylacetones of cerium, zirconium and thorium already mentioned, and salts with eightfold complex cations as

$$[Cu8NH_3][Pt(CNS)_4],$$

 $[Mg8H_2O][Th(NO_3)_6]$

or with an eightfold anion as in K₄[Mo(CN)₈], which gives rise to the free acid

H₄[Mo(CN)₈],6H₂O

There is, however, no evidence as to the spatial arrangement of associating units in these compounds. In a study of the alums which were formerly supposed to indicate a co-ordination number of 12, the tervalent metal is surrounded by 6 mols. of water, $[M^{III}6H_2O]M^I(SO_4)_26H_2O$. The arrangement of the remaining water molecules differs slightly according to the size of the univalent metallic ion (Lipson, Proc. Roy. Soc. 1935, 151, 347).

Reference should be made to compounds exhibiting odd co-ordination numbers (C.N.):

C.N. Three.—This number is displayed in a series of oxy- and thio-acids in which the principal valency of the central element may vary from three to five: horates [BO₃], carhonates [CO₃], thiocarhonates [CS₃], nitrates [NO₃], and sulphites [SO₃], meta acids of phosphorus, antimony, vanadium, columhium and tantalum [XO₃], chlorates [ClO₃], hromates [BrO₃] and iodates [IO₃].

C.N. Five.—Cupric halides co-ordinated with

5 mols. of ammonia, pyridine, etc., e.g.

[Cu5hase]X₂.

C.N. seven, of which series the analytically important K₂TaF₇ is the most outstanding example. It is noteworthy that tantalum also

gives a more regular type K₃TaF₈.

ELECTRONIO INTERPRETATION OF CO-ORDINA-

gives precision to the co-ordination theory since haltic sails such as $K_2[Co(NO_2)_2]$, and to diera a sumple explanation of the three manifestations of valency now recognized by chemists: (i) Electivation of valency in the polar or containable links as in the oppositely charged ions of platnam or the following series: $|P(NH_1|C)|_1$. sodium chloride, NaCl-, ammonium chloride NH. + CI or hexamminocohaltic chloride

(2) Covalency with non-polar non-ionisable links such as those which unite carbon to hydrogen in methane, nitrogen to hydrogen in ammonia, or carbon to natrogen in cyanogen. (3) The co ordinate link of Werner such as combines cobalt with ammonia in the foregoing cobaltammine.

The driving force discernible in these three types of linking is a tendency on the part of each of the combining atoms to assume the electronic configuration of the nearest mert gas. Of these gases helium has 2 electrons, hat each of the others has outer groups of 8 electrons—the

(1) Electrovalency (the Polar Link) -In forming acdium chloride, sodium gives its single outer electron to chlorine, and by this exchange the sodium ion acquires the electronic configuration of neon while the chlorine ion completes its

octet, thua resembling argon.

(2) Covalency (the Non-polar Link) —The two combining atoms share electrons as in methane, which contains 4 covalent links. Each of these 4 links includes 2 electrons, I contributed by carbon and I by hydrogen, so that in methane carbon has assumed the electronic structure of neon and each hydrogen has acquired the 2 electrons characteristic of helium. This type of non polar link to which each of two combining elements contributes 1 electron is typical of the non polar compounds of organic chemistry.

(3) Co ordination Covalency (the Co ordinate Link) -This mode of attachment differs from the foregoing type of co-valency or non-polar link in that both electrons are furnamed by one of the combining elementa (Sigdwick, JCS, 1923, 123, 725). This type of combination is

exemplified in the luteocohaltsmmine

[Co6NH,]CI,,

wherein the tervalent cohalt ion which differs electronically from krypton by 12 electrons completes ita quota of electrons by co ordinating with 6 mols, of ammonia. In this combination each nitrogen atom contributes 2 shared electrons, so that within the co-ordination aphere the cohalt has acquired the electronic configuration of krypton, whereas each nstregen retains the octet of neon.

By a similar process of reasoning it may be shown that this electronic equality holds in the case of purpurocobaltammine [CoCl5NHa]Cla and also generally among co ordination com-tion of the isomorphism of calcite and sodium pounds of tervalent cobalt. The fact that the intrate which were formerly formulated as acquirement of a complete octet by cobalt coincides with the symmetrical octahedral grouping of 6 associating units affords an explanation of the existence of so many welldefined members of this cohaltammine series respectively. But if the anions are [CO,] and

platinum of the following scries: [Pt6NH,]Cl; [PtCl5NH,]Cl;; and intermediate types to [PtCl₄2NH₂], [PtCl₅NH₃]K; and [PtCl₄]K_P

As regards co ordination compounds, chromium resembles cobalt, giving rise to numerous am-mines and complex salts, but in this instance the development of octahedral symmetry does not coincide exactly with the acquirement by the central metallic atom of an electronic steocture characteristic of Lrypton. Through. out the series there is a deficiency of 3 electrons. but, nevertheless, many individuals of the numerous group of co-ordinated chromic de rivatives show remarkable stability with similar phenomene of stereoisomerism and optically active enantiomerides.

Certain of the co ordination compounds of nickel and copper abow the opposits tendency in the combination of these bivalent metals with 6 associating units. For if each unit should contribute 2 electrons, then there would be an excess of electrons over the number required to endow the metallic radical with the electronic atructure of krypton Neverthelesa, such a complex salt as [Ni3dipy]Cla is sufficiently stable to be resolvable into optically active enantiomerides

This interpretation of co ordination as taking place with 2 electrons does not, however, exclude 1-electron covalent link. In the copper derivative (v. supra) of ethylenediamine is acetylacetone (Cuec) the cupric ion is still I electron short of the krypton structure which, however, is acquired when this compound co-

ordinates further with water or ethylenediamine hydrate to form respectively [H2OCuec] and [ecCuNH2C2H4.NH3]OH, the latter de-

hydrating to

[ecCuNH, C,H, NH,Cuec]

These addends are more loosely held than the original "ec," a circumstance which suggests the single link. In the nickel complex [Niec] the metal is still 2 electrons short of the krypton structure, and this unsaturation is revealed by the fermation of ammine and ethylenediamine compounds [2NH3Nlec] and

[C.H.(NH,),Nlec]

(Morgan and Main Smith, J.CS. 1925, 127, 2030 : 1926, 912),

NOTEWORTHY EXAMPLES OF CO-ORDIVATION PHENOMENA.

A Wider Basis for the Principle of Isomorphism. The older principle of integral valency led to structural formula which afforded no explans-

and accounts for the stability of complex co- [NO2] with a similar arrangement of oxygens

structure of these two compounds becomes explicable in terms of isomorphism. These ideas were developed by T. V. Barker (J.C.S. 1912, 101, 2484), who showed that the following substances were members of isomorphous series: K₂SO₄, K₂BeF₄, and (NMe₄)₂HgCl₄ (orthorhombic), KClO₄, BaSO₄, and KBF₄ (orthorhombic), xenotime YPO₄, zircon ZrSiO₄, and cassiterite Sn₂O₄ (tetragonal),

Cu[TiF₆]4H₂O; Cu[NbOF₅]4H₂O

and $CuWO_2F_4$, $4H_2O$ (monoclinic). The double salt K_2BeF_4 , $AI_2(SO_4)_3$, $24H_2O$ crystallises in cubic octahedra and is a true alum, as is also K_2ZnCl_4 , $Al_2(SO_4)_3$, $24H_2O$ (W. R. C. Curjel, Nature, 1929, 123, 206). In these substances the compounds $K_2[BF_4]$ and $K_2[ZnCl_4]$ are isomorphously equivalent to $K_2[SO_4]$.

Analytical Reagents.—The separation of copper from cadmium and of cobalt from nickel in qualitative analysis is based on the difference in stability of two pairs of double cyanides; (1) hydrogen sulphide decomposes the cadmicyanide, $K_2[Cd(CN)_4]$, but not the cuprocyanide, $K_3[Cu(CN)_4]$; (2) hypochlorites or hypobromites decompose the nickelocyanide

Na₂[Ni(CN)₄]

but not the cobalticyanide Na₃[Co(CN)₆]. The quantitative estimation of nickel is facilitated by the use of its insoluble complexes with dimethylglyoxime, a-benzildioxime, or dicyanodiamidine. 6-Naphthoquinoneoxime oneoxime dyes) gives a sparingly soluble cobalti-complex used in separating this metal from nickel. (see Chemical Analysis, vol. II, p. 612c).

Nitroso-\(\beta\)-phenylhydroxylamine (cupferron) furnishes insoluble red and white complexes with ferric and cupric salts respectively. The compound 8-hydroxyquinoline (oxine) has proved a useful analytical reagent for many metallic ions, since it yields insoluble products of the type indicated by the following general formula:

The reddish-violet colorations obtained by the action of alkaline copper solutions on biuret are complex salts $[(C_2O_2N_3H_2)_2Cu]K_2$,4 H_2O or $[C_2O_2N_3H_2Cu]K$,3 H_2O , in which this iminodiamide is reacting in its monoenolio form

HN:C(OH)·NH·CO·NH₂.

Mordant Dyes.—Included within the two main categories of mordant and acid mordant dyes are many substances of very varied chemical structure, but these otherwise diverse compounds have one feature in common, a chelate group capable of entering into co-ordination with the mctallic mordants generally employed in dyeing. The coloured effects produced on mordanted fibres are internal metallic complexes (insoluble lakes) arising from this combination of dyc and mordant.

round the central atom, then the rhombohedral or no colour. Alizarin yellow A and C are respectively benzoyl- and acetyl-pyrogallol (I), which, although but faintly coloured, furnish yellow aluminium lakes (II):

The quinoneoxime dyes are also simple examples of pale yellow to brown mordant or adjective dyes (chelate group HON: C-C=O) which furnish green and dark brown lakes on wool mordanted with iron and chromium respectively.

Alizarin (q.v.) and naphthazarin (alizarin black) supply further typical examples of such mordant dyes. The former (III) possesses 1 chelate group CO·C—C·OH, and the latter (IV) contains 2. When the hydrogens of the chelate groups are replaced by an appropriate metal aluminium, iron, chromium, etc.) a coloured lake is produced.

Treatment of III with pyroboric acetate reveals the presence of 1 chelate group (V), whereas with (IV) this reagent gives rise to a diboric ester containing 2 chelate groups (VI). Accordingly the quinone (IV) must be 5:8-dihydroxy-1:4-naphthaquinone and not the 5:6-dihydroxy analogue as formerly supposed (Dimroth and Ruck, Annalen, 1925, 446, 123.

An azo-group often becomes implicated in the chelate complex, as in the case of many hydroxy. In many cases the dyes themselves have little azo- and dihydroxyazo- dyes. Such colouring matters are aubstantive dyes on wool or silk | but their tinctorial effect is changed very considerably when the dye is converted into its form of fluorapatite, [Ca3(Ca3(PO4))]F, lakes with suitable mordants. Emochrome red (G. Shearer). The mineral portion of the bons B and Palatine chrome black 6B are examples of dispersions of vertebrates consists mainly of carbonato disportance of the producing give rise to deep red and black chrome comparing pressure to deep red and black chrome comparing places. In other instances the lake producing 1917, 111, 636; Hendricks, Jefferson, and chelate group is developed from the dye hy oxidation, as when the dyed fabric is developed with chromic acid. Diamond black PV, Carmoisin WS, and the chromotropes are typical examples of such dyes (J. Soc. Dyers and Col 1921, 37, 43; 1925, 41, 233) When these dihydroxyazo dyes contain aulphome groups it becomes possible to produce a soluble lake from dye and metallic mordant These soluble lakes which already contain the mner metallic complexes can be applied substantively to animal fibres and have the further advantage that they may be used in metallic vessels, since the lake is far less affected than the dye by the metal of the container A group of these soluble lakes is sold under the generic name of neolan dyes

The natural mordant dyes such as cochincal, lac, logwood and the redwoods also contain chelata groups capable of co ordinating with metallic mordants, and it is to the presence of such chelate groups (generally of the orthohydroxyketone type) that they one their time

torial properties.

Representative Minerals -The chemical conatitutions of many mineral species are most readily explicable in terms of co-ordination. The apatite group is a notable example, and it contains the following series chloroapatite, [Ca3{Ca,(PO,),}]Cl, pyromorphite,

mimetite, [Pb3{Pbs(AsOs)2)]Cl2, vanadinite, [Pb3(Pb,(VO,);)]Ci,

The spinels having the general formula Ham[RitO4] also form a notes orthy group of isomorphous minerals, including the following species. spinel, Al,[MgO,] hereynite, Ala[FeO4], chroniste, Cra[FeO4], magnesio-ferrite, Fe4[MgO4], gahnite, Ala[ZnO4], and magnetite, Fe [FeO] The naturally occurring solventes are also most renaily explanable at terms of the co-ordination theory. In these minerals silicon has a co ordination number of 4 as, for example, in the ion [SIO4].

Among copper minerals are the well known a ta camite

[Cu3(Cu(OH),)]CI,

and the corresponding sulphate, brochantite, [Cu3[Cu(OH)₄]]SO₄. Contrary to an opinion which has prevailed for a century the green patina developing on copper roofs is not base copper carbonate but hasfe copper sulphate, [Cux(Cu(OH),)]SO4, in which x sames from 1 to 3. In an artificial patina produced electro lytically, x=1, whereas in the natural patinas produced by exposure of more than 70 years a approximates closely to 3, so that the corrosion product is a co-ordination compound identical in composition with brochantite (Vernon and Whitby, J. Inst. Mctals, 1930, 44, 2, 389).

Substances of Physiological Significance - The fluorine in the enamel of teeth is present in the Mosley, Z. Krist. 1932, 81, 382), whereas the bone ash from fish and mammals contains as main constituent the complex phosphate

[Ca6|Ca1(PO4)2)(OH)2

(Morgulis, J Biol Chem. 1931, 93, 455). An important co-ordination compound of physiological significance in the animal kingdom

matter of blood Closely allied to this pigment is hæmocyanina, the blood colouring matter of crustaceans and molluses, but in this compound copper replaces iron. Another copper pigment of similar type is turacin, a reddish purple pro duct present in the feathers of the toursco

In the vegetable world, the chief co ordinated compound is chlorophyll, the green colouring matter of plants Two forms, a and β, have been recognised, and the structure of the former variety is indicated by II. The β form contains an aldehyde group replacing a methyl group, but

Merch [CH.], CHMe [CH.], CHMe [CH.], CMe CH CH.

both forms contain the 4- co ordinated magnesium atom.

Valency," by N. V. Sidgwick, Oxford University Press, 1927; "Annual Reports of the Chemical Society," 1930-1936.

G. T. M. and F. H. B.

COPAENE. The lævorotatory sesquiterpene hydrocarlion, copaene, $C_{1b}H_{24}$, b.p. 119–120°/10 mm., d^{15} 0.9077, $n_{\rm D}$ 1.48943, $[a]_{\rm D}$ –13.21°, occurs in African Copaiba oil (from Oxystigma Mannii) (Schimmel's Report, 1914, April, p. 48) and

References.—"Inorganic Chemistry, A Survey in oil of supa, Henderson, McNab and Robertson of Modern Developments," by Sir Gilbert Morassigned to it the structure (I) (J.C.S. 1926, gan and F. H. Burstall, Heffer and Sons, Cam-Stenzel (Ber. 1914, 47, 2555) as regards the bridge. On treatment with hydrogen chloride it yields cadinene dihydro-chloride (II) whilst on oxidation with ozone or potassium permanganate it gives a monobasic ketonic acid, copaene ketonic acid, C₁₅H₂₄O₃ (III), (semicarbazone, m.p. 221°), which on further oxidation with sodium hypobromite yields a dibasic acid copaene dicarboxylic acid, C₁₂H₁₈O₄ (IV).

Apart from cadinene dihydrochloride no

CHMe CH₂

$$H_2$$
C CH CH₂
 H_2 C CH CH₂
 C CMeCl 2HCl HC CH CMe

 C CHMe CH

 C CH

 C CHMe CH

 C CH

 C CHMe CH

 C CH

crystalline derivatives have been prepared from if so, the principal adulterants are gurjun copaene.

J. L. S. COPAIBA. Copaiba, or Balsam of Copaiba, is the oleo-resin obtained from various species of Copaifera, of which the most important is C. Langsdorfii. The oleo-resin is secreted in schizogenous ducts which eventually become lysogenous cavities in the trees, and these are opened by boring to allow the secretion to flow out. Copaiba is a thick, viscous liquid of a yellow to brown colour, often fluorescent. In commerce the balsam is sold on a description which indicates the place from which it is exported and which to a considerable extent covers the district in which it has been collected and, therefore, within reasonable limits, the species from which it has been obtained. The principal areas of its production are in Brazil

and the northern portion of South America.

Tho best known varieties are Maranham,
Maracaibo and Pará. The principal differences between the various balsams are in the proportions of essential oil and resin. Its use is cntirely confined to medicine, and the standards of the 1932 edition of the "British Pharmacopœia " are as follows. Its specific gravity should be from 0.960 to 0.995, and the acid value of the residue obtained by drying in a water bath should be from 120 to 160. The optical rotation of the essential oil obtained by distillation by steam or under reduced presence should be from -7° to -35° . The resin obtained by drying on a water bath should be not less than 50% and not more than 65%.

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balsam and the so-called African eopaiba.

Essential oil of copaiba (q.v.) is used to some extent in medicine, as is also the resin. For an exhaustive examination of a sample it is necessary to separate the essential oil and the resin and examine each separately. Tho essential oil is best distilled in a current of steam to avoid decomposition, which takes place even when the balsam is distilled in vacuo. The table at the top of p. 338 will show the figures for a number of typical authentic samples, but slight variations may be expected in a substance derived from several species under varied conditions. These values are merely typical, variations from them may be expected, and definite limits cannot be given.

For the detection of gurjun balsam oil, tho determination of the optical rotation of the essential oil is important. Gurjun oil has a rotation which is rarely below -70° and often reaches -130°, so that a higher rotation than -35° in the essential oil renders a sample very suspicious. If 3 or 4 drops of the essential oil are dissolved in 3 c.c. of glacial acetic acid and the mixture is carefully poured on to the surface of concentrated sulphuric acid, a dark violet colour is developed in the acetic acid solution if 5% of gurjun oil be present.
The resin of copaiba consists for the most part

of bodies of an acid nature. According to Schweitzer (Pogg. Ann. 1829, 17, 784; 1831, 21, 172) one principal constituent is a crystalline ot more than 65%.

At present copaiba is seldom adulterated, but acid was assigned. Its formula is given as

	Marsaham	Maracalbo.	Cartagena	Pará,	Bahls.
Sp gr. of balsam at 15 5° Acid value of balsam Refractive index of bal-	0 990 70–80	0 980-0 995 85-90	0 950-0 970 80-80	0 920 30–55	0 975 80
Retractive meet of parama (20°) Essential oil % Sp.gr. of oil at 15.5° Refractive index of oil at 20° Optical rotation of oil at 20° Acid value of resin Ester value of resin		1·515 45-48 0 900 1·4980 10° 80-90 50	1.510 42 0.904 1.4960 -20° 130 45	1 508 75 0 900 1-498 -25° 70 85	1 506 44 0-905 1 495 -18° 75

C₂₀H₂₀O_F. Numerous other acids have been described, but as little other then combustion results and empirical formula have been quoted, it is not necessary to reproduce them here (see Arch. Pharm. 1901, 235, 648).

Species of Co- paifers	reticulata	guyanensıs	multijuga	officensles	Martin, v Figida	coriacea	Martsi	glycy- carpa
Place of col- lection	Tspajóz	Rio Negro	Tapajós	Rio Negro	Pará	Bahla	Santarem	Maués (Amaz)
Sp gr (15°/4°) Optical rota- tica (100	0 94-1 004	0 93-0 975	0 92-0 955	0 95-0 998	0 93-0 95	0 925–0 965	0 888 -0-91 2	0 958-0 si
mm tube, at 20°)	-85* 40°/ -37* 10°	-28* 30"	Not det	ermined	-14* 45%	- 22° 30'	Not dete	rmined
Acid no .	74-81	68-71	81-86	61-63	-19° 20°	-28° 15° 58-82	48-67	65-82
Bapoaifica- tion us. Solubility la 90% alco-	89-91	92	77-82	62-70	68-88	Not det	44-62	72-11
hol: vols,	6	8.5	6	45	12	10	5.5	12
Essential oll,	26-31 5	28 5	48 5-55	55-58	60-63	55-62 5	38-52	32-38
8p gr of oll,	0 935/0 938	0 910/0 912	0 895	0-915/0 918	0 926/0 929	0 911/0 914	0 922/3 928	0 925/0 9

119); end von Soden (Chem. Ztg. 1909, 33, It will be convenient to describe here the

so called African copaths. This body is also known as Illuran balsam. Its botanical origin is nacertain. It has been ascribed to Oxystiqua Mannii Harms, but is more probably the product of Daniellio thurifera. It closely resembles true copaiba in characters, the principal physical difference being that its essential oil is dextrorotatory. The characters of the balsam are as follows:

Specific gravity . . 0 985-0 998 Acid value . 55-60 8-10 Ester value

Illuric acid, C20H24O2, melting at 1280, is stated to be the chief of the resin acids (Tachirch) (absence of oil of African copasha). The oil " and Keto, Arch. Pharm. 1901, 239, 548).

COPAIBA, ESSENTIAL OIL

See also Balsam of copaths, "Allen's Com-mercial Organic Analysis," 5th eds. Vol. IV. Copat/en (Fam. Leguminose), which are large p. 307; Deussen and Eger (Chem. 24; 1912, 33, trees found chefly in Braul, Veaczuela, Gunza 561); Cocking (Chem. and Druggest, 1916, 77, and Colembia. The oleo ream is a scertifica which is obtained by boring the tree and collecting the exudation. It is imported from Maran-ham, Maracaibo, Pará, Cartagena, and Cayenne. The Para variety yields 85-80% of oil, the other varieties 35-559

Composition -This oil consists mainly of the sesquiterpeace, \$\beta\$ caryophyllene and cadinene.

Characters .- It is a colourless or pale yellow oil. ap gr. 0 895-0 908, opt. rot. -7° to -35°, rel ind. 1 495-1-500. A solution of 1 ml. of oil m 5 ml. of glacial acetic acid ahould not develop more than a faint reddish or purple coloration of the addition of 4 drops of nitrio acid (sheence of gurjun balsam). When distilled under reduced pressure the first 10% fraction has an optical rotation lower than that of the original oil used in medicine as e urinary astiseptic.

COPAL,-This varnish resin is, from The oil distilled from the oleo resun or balsam commercial point of view, of very great in-

portance. It will he convenient to deal with, its numerous varieties, from a geographical point of view. The principal of the copals are as follows:

(1) East African copals.(2) West African copals.(3) Kauri gum or New Zealand copal. (4) East Indian and Manila copals.

(5) South American copals.

(1) The trees yielding East African copal (Zanzihar, Lindi and Tanganyika) are Trachylobium verrucosum (Fam. Legamimosæ), Hymenæa (Madagascar) = Trachylobium ververrucosa rucosum and Copaifera conjugata Milne-Redhead (Inhamhane).

The resin ducts are near the surface and no injury to the tree is necessary. Native collectors recognise three varieties of copal: (a) that ohtained direct from the trees; (b) that which has dropped from still growing trees and collects in the soil, where it has hardened, known as "semi-fossil" copal; and (c) that found in the soil, which has been dropped from trees no longer existing, known as fossil resin. Some of the fossil resin has characteristic rough markings, and is known as "goose skin" copal. This is probably due to uneven contraction of the various layers during drying.

The following analyses of East African copalknown generally as Zanzibar, although the greater part is produced on the mainland—are due to the Belgian Ministry of Colonies (Revue générale de la Colonie belge, 1921,

552).

	Sp. gr.	Mois- ture. %	Ash. %	soluble in alcohol.	Acid value.	Sap. value.
1 2 3 4 5 6 7 8 9	1.062 1.071 1.062 1.064 1.067 1.061 1.074 1.053 1.057	0.81 0.87 0.80 1.15 0.89 0.81 0.66 0.59	0.04 0.04 0.04 0.05 0.02 0.06 0.08 0.04 0.01	53·16 57·88 48·20 52·22 54·53 51·72 93·74 93·18 81·39	83·1 86·7 87·6 100·7 100·1 95·9 153·9 166·9 142·7	83·1 89·5 89·1 100·7 100·1 96·0 165·1 166·9 151·1

Similar analyses of Madagascar copal (l.c.) gave the following results:

!	Sp. gr.	gr. Moisture. %		soluble in alcohol.	Acid value.	Sap. value.
1 2 3	1.062	0·52	0·014	76 05	140·3	155·7
	1.060	1·09	0·40	57 31	111·08	140·3
	1.052	1·05	0·27	49-28	100·8	111·8

The constituents of Zanzihar copal are not known with any degree of certainty. Resenes and acids with very high molecular weights have heen quoted, hut until a series of well-defined derivatives has been prepared, the individuality of these bodies cannot he relied on. Reference may he made to the papers of Tschirch and his fellow workers (Arch. Pharm. 1896, 234, 552; 1897, 235, 92; 1901, 239, 145, 298; 1902, 240,

202, 478; 1908, 246, 293).
(2) Of West African copals, Congo copal is by far the most important. The greater part of the exports consists of fossil resin. Sierra Leone copal is obtained from living trees. Various other copals or so-called copals are produced in other West African districts. The copal from the Belgian Congo is derived chiefly from Copaifera Demeusei Harms, and to a smaller extent from other Copaifera species. A large number of samples have been examined by the Belgian Ministry of Colonies, which have been

Specific gravity . 1.033-1.065 Moisture % . 0.26-1.000.017-0.06 38.82-70.53 85-4-143-7 Acid value Saponification value 91.1-149.5

collected in numerous districts. These samples

gave results within the following limits:

For Tschirch's views on the composition of Congo copal, see Tschirch and Engel, Arch. Pharm. 1908, 246, 293.

Hellinckx gives figures for Congo copal which are of value, since the samples are all graded as to colour:

No. Grade.		Melting-	point °C.		4		Iodine value.	
	Appearance.	Lower.	Upper.	Sp gr.	Acid value.	Sapon. value.		
1 2 3 4 5 6 7 8 9 10 11 12	White ", ", Dark ", Amher	Pale brown, white strictions Brownish, opaque	120 — 110 110 108 135 132 132 135 90 126 130	178 	1.0509 1.0585 1.0482 '1.0459 	100 130 117 104 — 98 114 114 84 85 90	120 136 132 130 — 128 150 150 150 150 130	104 82 73 90 — 70 74 54 59 96 14
13 14	,,	Brown, chalky appearance . Yellow opaque	125 95	195 145	1.0550	79 42	121 59	110 72

It will be seen that the analytical examination always apply to kauri copal. The following of copal must be regarded as quite aubsidiary table fairly represents the average hardness of to the expert knowledge gained by those copals taken in their order, with Zanzibar copal accustomed to handle specimens of known at the head of the list ;

(3) Kauri gum or New Zealand copal is the product of Agathis australis (Fam. Araucariacea), the fossil "gum" being dng from the soil.

The tree is said to live for over 3,000 years Kauri "gum" varies greatly in quality, as some is found in fine, clear, bold pieces, some as small fragments, and some adherent to old tree stumps, from which it has to be extracted.

This resin has the following characters:

2-5
0-02
60-80
84-130
2-20
100-125

Tachirch and his fellow workers have, as with most of the copals, given formulæ for various acids isolated from this resin, but the most recent work on the subject is that of Ruzicka and Hosking (Annalen, 1929, 469, 147; Helv Chim. Acta, 1930, 13, 1402, sond. 1931, Chim. Acts, 1930, 10, 1400, 1001.

14, 203 They isolated a crystallina acid,

C₂₀H₃₀O₄₀, which they termed agathohicarboxylic acid (q t.) It is present in both New Zealand and Manila copals. An ethersolubla amorphous resin acid from Manila copal on dehydrogenation with selenium gave 1:2:5trimethylnaphthalene, a hydrocarbon, C1,H4. and retene, whilst agathre acid yielded pimanthrens but no retens

(4) The East Indian and Manila copals, which include the copals from the islands of the South Pacific from Sumatra and the Malay Peninsula on the west to Celebes and the Philippins Islands on the east, are derived mostly from the living tree Agathis alba. These copals have

tha following characters

Acid value (of pure resin) . 120-150 Saponification value (of pure resin) 165-190 Iodine value . .

For the older viewage to chemical composition. see Tschirch and Koch (Arch. Pharm. 1902. 240, (3), 202) and Richmond (Philippine J. Sci. 1910, 5, 177).

Ruzicka, Steiger and Schinz (Helv. Chim. Acts, 1926, 9, 962) have examined Manila copal and found it to consist of 80% of acids as followa:

 (1) A monobasic acid, C₁₄H₂₄O₂ nr C₁₈H₂₂O₃
 (u) A dibasic acid, C₂₃H₂₄O₄, apparently identical with Pichmond's acid, 233H34O4

(ni) A dibasic acid. C40H20O2

(5) The South American copals are prebably derived from species of Hymenau, but are not of great commercial importance.

In general properties the copals vary in colour from white to yellow, red, brown and brownish-black; their softening and meltingpoints and their specific gravities also vary. An extellent criterion of the value of copal is its amyl alcohol or ethyl acetate, and regained hardness. Generally speaking the harder the on cooling or on reprecipitation of the co pigment copal the greater its value, but this does not into the aqueous layer.

(1) Zanzihar. (8) White Benguela. (2) Mozambique. (9) Cameroon. (3) Lindi. (10) Congo. (4) Red Angola. (11) Manila. (5) Pebble Copal. (12) White Angola. (13) Kauri.

(6) Sterra Leone, old. (7) Yellow Benguela. (14) Sierra Leone, new.

Copal is partially soluble in many organic solventa, for example, in alcohol, especially affer it has been left to swell up in other But to convert ropals into a completely soluble form it is necessary to heat them, sometimes even to the point of destructive distillation at 400°C., when the resin becomes soluble in turpentine or linseed oil.

Many analytical values have been published for copal in general. The following papers should be consulted . R. Williams (Chem. News, 1888, 58, 224); Lippert and Reissiger (Z. angew. Chem. 1900, 13, 1047); Schmidt and Erban (Z. angew. Chem 1889, 1, 35); Worstall (J. Amer Chem. Soc 1903, 25, 860); Hellinckr (Dissertation, Louvain University, 1935).

Iodine Value.—The most full examination of the iodine value is that of Worstell (I.c.) and for Congo copal, Hellinckx (lc.). gives the following values hased on the Hubl method.

_					Minimum
Турс				Maximum	MARKETTE
Kaun				. 170	74
plands				. 148	104
Pontian				. 142	119
Zanziba				. 123	115
Mozami					136
Madaga					26
West A				. 143	122
Sierra L		8		. 105	102
Brazilia	n			. 134	123
					E. J. P.

COPALIN. Highgate resin. A fossil resin found in the blue clay of Highgate Hill.

The influence of tannia CO-PIGMENTS. in increasing the blue tone of anthocyanin flower pigments when they are in solution was 5rst observed by Willstatter and Zollinger (Annalen, 1916, 412, 212). The phenomeno was further studied by G. and R. Robinson (Bjochem. J. 1931, 25, 1687; 1932, 28, 1647). who discovered the importance and significance of nnt only tannins but of other colourless enbstances in flower colour variation. They termed all these compounds which turn authocyanins blue "co-pigments." Typical copigment effects are seen in magenta and rosepank Chanese primulas as rompared with red and salmon ones, and in the purple inner petals of the fuchsia.

A co-pigment effect is visible in criro as well as in tire, but is lost at boiling point or on extraction with an immiscible solvent, such

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and exist for all types of anthocyanins, but with a few outstanding exceptions pelargonidin derivatives, especially 3-biosides, are not found to be greatly affected by co-pigments. The chief natural co-pigments appear to be the tannins and antboxantbin (flavone and flavonol) glycosides. Experimenting with synthetic compounds, G. and R. Robinson found that while 2liydroxyxanthone, papaverine and quinaldine act as strong co-pigments on specific anthocyanins, 4-bydroxyxanthone, laudanosine and quinoline have very little action. They suggest that the phenomenon has little or nothing to do with salt formation, since it occurs in the presence of a large excess of mineral acid; but is evidently the result of the formation of weak additive complexes which are dissociated at an elevated temperature or by the action of solvents.

Co-pigments not only modify the colour of acid anthocyanin extracts, but they also have specific effects upon distribution numbers and upon the colour reactions with ferric chloride, sodium carbonate or sodium acetate.

Robinson, Robinson and Roberts (Biochem. J. 1934, 28, 1712), using narcotine and papaverine as co-pigments for oenin and chrysanthemin, found that the extent of the depression of the normal distribution number (the partition valuo between amyl alcohol and aqueous acid) was correlated with the extent of the co-pigment effect. Papaverine was in both cases much more effective than narcotine. On plotting the log. curve for the distribution of oenin or chrysanthemin hetween amyl alcohol and water at various concentrations a slope of 1 instead of 2 was obtained when excess papaverine was present. Since it had already been deduced by Levy and Rohinson (J.C.S. 1931, 2720) that in aqueous solution anthocyanins were associated as double molecules, while they were single in the bluer amyl alcohol layer, these results with co-pigmented extracts point to the breaking up of the double molecules by a combination between alkaloid and anthocyanin, thus giving the distribution and blueing effect compatible with single molecules. When compared over a large range of p_R values, copigmented anthocyanins are always found to be bluer than the normal pigment (cf. Buxton, J. Genetics, 1932).

The researches (Robinson et al.) upon the marked specificity of co-pigment action were followed by Lawrence's observations (Nature, 1932, 129, 834) upon co-pigmentation in Dahlia, and upon the remarkable variation in the specific effects of various crude ivory antho-xanthin extracts upon a number of natural authocyanins (see Scott Moncrieff, J. Genetics, 1936, 32, 117). The effect of these ivory pigments upon malvidin 3-monoglycoside was extreme; with cyanin the effect was strong but not so marked, while pelargonin was only slightly modified. Moreover, each anthocyanin was modified in a different degree by each of the anthoxanthin extracts.

Since the production of anthoxanthins and of anthocyanins is inversely correlated, a co-

These compounds are specific in their action | paler in colour than the corresponding red or purple unco-pigmented variety (Lawrence and Scott-Moncrieff, J. Genetics, 1935, 30, 155).

R. S.-M.

COPPER (Fr. Cuivre; Ger. Kupfer). Sym. Cu, at. wt. 63.57, at. no. 29, isotopes 63, 65.

OCCURRENCE.—Copper occurs in the native state in many localities and is widely distributed as compounds of the metal, the following being the most important minerals from which it is extracted: native copper, sometimes associated with silver and hismuth; oxides, cuprite Cu₂O, tenorite or melaconite CuO; sulphides, chalcopyrite or copper pyrites Cu₂S, Fe₂S₃, chalcocite or copper glance Cu₂S, covellite CuS, bornite or erubescite Cu₅FeS₄; double sulpbides with antimony or arsenic, tetrahedrite 4Cu₂S,Sb₂S₃, tennantite4Cu2S,As2S3,famatinite3Cu2S,Sb2S5, enargite 3Cu2S,As2S5, bournonitc,

$3(PbCu_2)S,Sb_2S_3;$

sulphate, chalcanthite \mathbf{or} copper vitriol CuSO4,5H2O; carbonates, malachiteCuCO₃,Cu(OH)₂, azurite, 2CuCO₃,Cu(OH)₂; silicates, chrysocolla CuO,SiO₂,2H₂O, dioptase CuO,SiO₂,H₂O; chloride, atacamite,

CuCl,,3Cu(OH),.

The importance of the several species of ores in the above list varies very much with the locality. Thus, while native copper occurs to a considerable extent in the Lake Superior district of North America and at Monte Catini, Italy, and is worked for the metal, and atacamite occurs and is worked at Los Remolinos and tho Atacama Desert, South America, these minerals would in other localities be looked upon as rather rare, and as occurring only in the zone of weathering of copper lodes. So also in the case of malachite, this mineral, as an ore of copper, has heen seldom worked except in small quantities in the upper and oxidised portions of some copper lodes, yet in the case of the once celebrated Burra Burra mine in South Australia and at Nishni-Tagilsk in Siberia it formed the bulk of the ore produced, and very largo quantities have been obtained from these districts. The most remarkable deposit of malachite is that of the Katanga region of the Belgian Congo and the adjacent part of Northern Rhodesia; this is a most important field for copper and produced over a quarter of a million tons of the metal in 1936; the minerals are malachite, azurite, chrysocolla, melaconite, chalcocite, together with, at some mines, chalcopyrite and bornite; chalcopyrite and other sulphides are encountered at depth.

Chalcopyrite is the principal commercial source of copper and occurs in a number of ways, the most important of which are hydrothermal veins as in California, Montana, Arizona, Alaska, and Canada, pyrometasomatic deposits in which the chalcopyrite occurs with other sulphides at or near the contact between bodies of intrusive rock and limestone as Clifton-Morenci and Bisbec, Arizona; Bingham, Utah; Alaska, Canada, Australia, Japan, and pigmented magenta or blue flower is necessarily Korea. Chalcopyrite associated with bornite

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grains in a shale and are of sedimentary origin ments.

deposited at the same time as the shale.

States, Chil, and the Katanga area of the of the smelting process is to obtain the copper Belgian Congo and Rhodesia each produce about one-fifth. Copper ores frequently carry a low percentage of copper, thus some ores worked at a profit carry only 0.5% of the metal, and the average content is probably about 20%. The ore reserves of the Nchanga Consolidated Copper Mines of Northern Rhodesia are estimated at about 128 million tons having a copper content of 4 66%. The Nchanga West portion of this total is 41 million tons which includes 51 million tons containing 17% copper. The importance of the Northern Rhodesia deposits can be realised from the fact that in 1936 the ore the silica originally present in the ore, comreserves were reported as 500 milhon tons, with an average content of 411% copper (Parker and Gray, Trans. Inst. Min. Met. 1936, with earthy hases also present or with lime added 45, 330)

EXTRACTION.

Generally speaking, in all copper ore deposits a certain amount of oxidised material is found in the portions near the surface, but as depth in the decomposition of this matte in such a is attained and almost invariably at or about manner as to eliminate the iron and sulphur the level at which water is reached, the copperbearing minerals change and the oxidised minerals give place to sulphide minerals, and these finally to chalcopyrate intimately mixed with varying proportions of pyrite. Chalcopyrite thus diluted down with pyrite, pyrotite, etc., forms the ore from which the greater proportion of the world's copper supply is derived, and the hulk of the material treated at smelting works generally. Such ore sometimes contains appreciable quantities of gold and silver. The silver is frequently carried by intermixed fahl ore, but also in many instances by the pyritic mixture in which distinct aliverbearing minerals are not observable. The ores as mined generally contain only small percentages of copper, frequently not more than 1%, and are submitted to some form of concentration, the most important being flotation processes hy means of which a concentrate is obtained much richer in copper and more suitable for treatment and a tailing containing very little copper, which can be discarded.

The actual extraction of the metal and production of copper in the metallic state is carried out by one of two distinct types of process; one of these, hy far the most important because of its very general application, treats the ore by smelting and is known as the dry method. The other-of only limited application-treats the material with solvents. thus obtaining the copper in solutions, from which it is recovered by suitable means, these

processes are known as wet methods.

taken into account in deciding whether a dry or being the concentration of the copper in the a wet method is best for any given ore is the matte, the chief considerations kept in view nature of the ore itself, but other considerations are the production of a matte as rich in copper

occurs in the Kupferschiefer at Manafeld, includes the availability of suitable fuel supplies, Germany, where the copper minerals occur as water supply, suitable is bour, and other require-

Dry Methods .- These methods are autable The world's production of copper is about 11 for ores in which the copper is present as million tone per annum, of which the United sulphide, and the object of the first stage in a concentrated form in the first product of smelting. For this purpose the ore is smelted under exidising conditions in such a way as to produce as the mam product a "matte" regulus" which consists of cuprous sulphide Cu.S. and aron sulphide, FeS, in varying proportions, together with sulphides of certain other metals which may be present in the ore. Practically the whole of the copper present in the ore thus passes into the matte and a second product containing only traces of copper, alag, is also formed, The slag consists of hined with oxide of iron formed by the oxidation of part of the iron sulphide present together to the charge. The alag is thus mainly composed of iron silicate, and owing to its lower specific gravity separates easily from the matte and is removed to the slag dump.

Having thus concentrated the copper in the matte, the second stage of the process consuts and, as far as possible, other metallic elements which may be present and to yield the copper in the metallic state. For this purpose complete oxidation of the iron sulphide present and the combination of the oxide of iron formed with ashea to form a slag is carried out, together with partiel oxidation of the cuprous sulphids and subsequent reaction of copper sulphids and oxide yielding metallic copper. This is not sufficiently pure for the market as it may contain small quantities of impurities originally present in the ore and it may also contain gold and silver worth recovering. This impure copper is commonly known as blister copper, from the fact that large blusters are found on the surface of the piga of copper, formed by the evolution of sulphur dioxide from the metal

during its solidification. The final process therefore consists in the refining of the copper produced as above, or in removing as far as practicable the small quantities of such elemente as may still be present which would either injuriously affect the metal for manufacturing purposes or be of sufficient value to be worth separating.

The treatment of the ore involves, therefore,

three distinct operations: L Smelting the ore with the production of

matte and slag 2. Treatment of the matte for the production of crude copper.

3. Refining the crude copper.

1. SMELTING THE ORE WITH THE PRODUCTION The main consideration which has to be OF MATTE AND SLAG. The object of this process include the locality of the ore deposit, and this and a slag as poor in copper as is practically

of matte for subsequent treatment having heen decided upon, the smelter regulates this by suitably proportioning the sulphur in the charge to he treated so that there shall he sufficient to form Cu2S with all the copper present and a further quantity to form FeS with some of the iron present so that the matte produced shall contain the required percentage of copper. From these considerations it will he seen that the factor which regulates the grade of the matte is the proportion of sulphur to copper present in the ore mixture to he smelted. When smelting purely pyritie ores or concentrates it frequently happens that the quantity of sulphur present is considerably in excess of requirements, and if smelted direct would result in a low grade of matte, so that the production of the most suitable grade necessitates the removal of this excess of sulphur hy a preliminary operation known as " calcining" or more properly "roasting."

This process consists in subjecting the ore to a dull red heat under oxidising conditions so that the sulphur combines with oxygen derived from air admitted and in contact with the ore with the production of sulphur dioxide, which is generally used for the manufacture of sulphuric acid, while oxide of iron and oxide of copper to some extent are formed at the same time.

The method of carrying out this roasting process depends upon the nature of the ore and also upon the locality; thus it may be roasted:

(a) In the rough or lump form—in pieces from

1-in. cuhe upwards; or

(b) In the finely divided state or as concentrates ohtained hy preliminary mechanical washing of the ore or hy flotation.

Under (a): Copper ores were formerly roasted in heaps in the open air by starting the comhustion hy means of wood, the comhustion heing maintained hy the sulphur in the ore. It is evident that there must be objections to this method owing to the contamination of the atmosphere with sulphur dioxide, and there are few localities where such a method would now he tolerated. As an improvement on heap roasting, stalls and kilns have previously also heen used; these consisted of series of chambers huilt back to hack, suitable flues being arranged to carry the fumes to a chimney stack. An improvement on these methods consisted of the use of "burners" or closed chambers similar to those used for burning pyrites in sulphuric acid works, the sulphur dioxide heing used for the manufacture of sulphuric acid.

Under (b): Finely divided copper ores were formerly roasted in long-hedded reverberatory furnaces to which a firehox was attached at one end and an exit for the gases at the other end. The ore was dropped from a hopper through an opening in the roof on to the bed at the fluc end of the furnace, where it was spread out hy suitable tools worked hy hand. The conditions in the furnace were intensly

and economically possible. The desired grade | being discharged through openings in the hed into vaults or chambers placed below. The labour required on these furnaces was very heavy and the output small, so that they have now been displaced by other methods of roasting, including those in which the ore is stirred, moved forward, and discharged hy mechanical

means operated hy power.
Several forms of straight-hearth mechanical furnaces have been used similar in principle to the hand calciner. For example, one such calciner consists of a flat hed, 100 ft. or more in length, covered with an arch. At the cool end is the inlet for the ore and at the opposite end is the outlet. At the hot end also, hut on the side of the furnace, is a firehox with its inlet to the furnace constructed in the roof of the latter. Other similar firehoxes are placed along the length of the furnace, and at the cool end is the flue for carrying off the gases. A channel is constructed in the side walls of the furnace in which rails are laid and suitable carriages with wheels resting on the rails stretch across the furnace, to these ploughs are attached which, passing through the furnace, turn over and mix the ore and at the same time carry it forward. The carriages and ploughs are moved by a pair of endless chains which cause them to travel through the furnace in one direction and to return in the other direction under the furnace bed. Each time the carriage enters the furnace it takes in with it a portion of ore or concentrate to he roasted and each time it leaves the furnace it carries out a

similar portion of roasted product.

The McDougall furnace (Fig. 1) is typical of the class of roaster largely used at the present time. This consists of a series of flat, circular, superimposed firebrick hearths enclosed in a hrick-lined cylinder of steel plate. An iron shaft passes centrally up the cylinder through openings in the several hearths and is caused to revolve hy suitable gearing attached to one end. This shaft is cooled hy water or air. Arms are attached to this shaft above each hed. These arms extend from the shaft to the periphery of the circular heds and have attached to them a series of ploughs set at a slight angle to the arm. The ore is fed from a hopper on to the top hed near the outside and as the arms revolve they turn the ore over and gradually move it from the outside to the centre of the hed. At this point is an opening through which the ore falls on to the second hed. On this hed the ploughs are so set on the arms that the ore travels in the reverse direction and is drawn to the outside of the hed. Here it discharges on to the third hed and so on until it passes out at the hottom of the furnace. Arrangements are made for the charging of this hot roasted material into reverberatory furnaces with as little fall in temperature as possible, leading to economy in fuel. The openings through which the ore falls from one hed to the other serve as flues through which the gases pass upwards through the furnace, finally escaping through a flue above the top hed to the sulphuric oxidising and the charge was slowly moved acid plant. Attached to the lowest had is a from the flue end to the firchox and hy the firebox hy which the furnace is heated up in workmen, losing sulphur on its way, and finally order to start the roasting; after once being

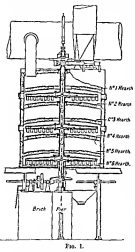
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started, little or no fuel is required, as the most of the material is at the present time oxidation of the ore provides sufficient heat owing to the development of concentrating throughout the apparatus; a partial removal of methods and especially of flotation processes,

sulphur is all that is required,

Another method of removing the excess sulphur which has been used is known as " pot roasting" or "blast roasting." This method has been applied to concentrates and fines and, fornace amelting, and for this blast furnaces are while it eliminates the sulphur, the heat attained is at the same time sufficiently intense to cause the ore to partially melt or amter together, forming a mass which requires to be broken up but is suitable for blast-furnsco trestment.



The process is carried out in a similar manner to that used in the treatment of lead and zinc ores but is little used in the metallurgy of

After the sulphur in the ore mixture has been reduced to the requisite amount it is ready to be smelted. For this operation one of two distinctly different types of furnace is used, material placed in position under the spout. These are: (1) blast furnace; (u) reverberatory furnace.

it is generally unsuitable for blast furnace treatment, while it is emmently suitable for the reverberatory furnace. On the other hand. rough ore is unsuitable for reverberatory used Certain ores which carry gold and silver in the gangue minerals and ores which contain both sulphide and oxidised minerals of copper are best treated in blast furnaces. During the smelting of the former, the precious metals will pass into the matte formed and will be carried into the metallic copper obtained. from which they are easily recovered during electrolytic refining

(i) Blast Furnare Smelting -The furnace used is constructed with water cooled walls of steel plate, each section of which is termed a jacket and is generally rectangular in shape. As the working of this type of furnace is dependent upon an air blast projected into the interior, the width of the furnace in limited by the strength of the blast which is required to penetrate the charge from the sides to the centre of the furnace, while the length of the furnace is only limited by the capacity required. The increase in length is obtained by clamping together two or more of the jackets on each aide, thus extending the longer axis of the furnace

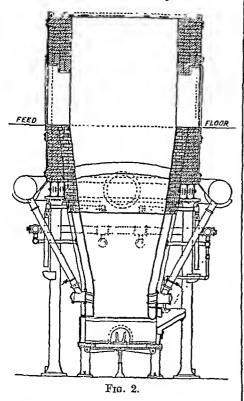
The jackets at and on a thick cast iron plate carried on iron supports. In some cases, as in Fig 2, these jackets are only used in the lower part of the furnace, the upper part being constructed of brick, but the brickwork is new generally replaced with a second tier of jackets reaching up to the feed floor. Above the feed floor the furnace is continued in the form of a covered-in brick chamber with lifting doors for charging and an outlet for the waste gases. The charge consists of a proper mixture of ore-either raw, raw and calcined, or partially calcined—flux, either limestone or allicious material, and the necessary amount of coke required to form a suitable smelting mixture On the lower floor the furnace is provided with spouts through which the molten products of the furnace flow. The aide jackets of the furnace are provided with circular openings forming a borizontal line of openings on each ade of the furnace through which the tuyeres pass for the admission of the blast. The blast at a suitable pressure is generated by a blowing engine and a plentiful supply of water is circulated through all the jackets, the overflow pipes carrying away the hot water.

The products of the smelting operation carried out in this furnace flow out through a spout into the settler. This settler consists of a tank, constructed of steel plates haed with refractory It may be 10 ft, or more in diameter and in provided with a slag apout near the top and a The choice between these methods of smelting tap hole near the bottom. In this vessel s depends on a variety of circumstances, all of separation of the matte and slag takes place; which must have proper consideration, but the the matte having the higher specific gravity falls following are the chief points, viz.; if the ore to the bottom and the lighter alag rises to the to be smelted is in the state of concentrates as top, whence it overflows through the slag notch

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into suitable pots provided for its removal without drying, roasting, or sintering. to the slag dump. From time to time the matte is tapped off through the tap hole at the bottom of the settler for further treatment. The working of this type of furnace proceeds continuously, fresh ore mixture and coke being charged at short intervals while a constant flow of matte and slag passes out through the spout into the settler.

Pyritic Smelting.—This method of working in the blast furnace secures in one operation that which in the preceding description is attained in two processes carried out in separate furnaces. In pyritic smelting the high temperature necessary to carry on the smelting is derived from the oxidation of the sulphur and iron



(chiefly the latter) contained in the ore itself. While this method of working without coke as a fuel is of the highest importance from an economical point of view, it is of limited application owing to the fact that comparatively few mines produce ore which, as regards its physical and chemical constitution, is suitable for this method of treatment. In working the process it is found advantageous to use from 1 to 3% coke instead of the 10 to 12% used in the ordinary blast-furnace method of smelting roasted ore. A small amount of coke used alters the process to a partial pyritic smelting, and is found to be desirable in order to keep the charge open.

In the Mt. Lyell smelter practice the flotation concentrate carrying 26% copper and 12%

The blast-furnace treatment of this fine material has necessitated the introduction of an efficient dust-collecting system in which 30 tons of dust per day, containing 25% copper, are obtained. This dust is incorporated with the concentrate pulp prior to filtration, and so is returned to the blast furnace (J. N. Greenwood, Proc.

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Austral. Inst. Min. Met. 1936, No. 103, 225).
(ii) Reverberatory-Furnace Smelling.—The furnace used in this process is essentially different both in construction and in method of working from the blast furnace. Instead of the fuelcoke-being burnt in contact with the materials to be smelted and the combustion being carried on by means of an air blast, the material to be smelted occupies one compartment of the furnace while the fuel, if lump coal be used, is burnt in another. The draught required for the combustion of the fuel is quite moderate and is obtained by a chimney of suitable height and area. The use of lump coal firing has given way almost entirely to the use of pulverised coal or oil fuel which is introduced through suitable burners. The working bed of the furnace in plan is a parallelogram. At one end and separated from it by a low partition is the firebox or space for necessary burners. At the opposite end is the outlet flue connecting with the chimney. A brick arch extends from the firebox end to the flue and covers the furnace from side wall to side wall. The side walls of the furnace contain a number of openings used in working the process for spreading the ore charge, drawing off the slag, etc., and one or more tap holes are provided for tapping the matte (Fig. 3).

The furnace walls and roof are constructed of highly refractory bricks with an outside casing of ordinary bricks, and the whole is bound together with steel girders placed vertically and tied together by rods. The charging of this furnace takes place from hoppers situated above openings in the roof near the side walls and extending half the length of the furnace from the firing end, and hot, partially roasted concentrate is often used. The charge melts down and reactions take place between oxide of copper and sulphide of iron present whereby the whole of the copper is converted into sulphide, according to the equation:

Cu₂O+FeS=Cu₂S+FeO.

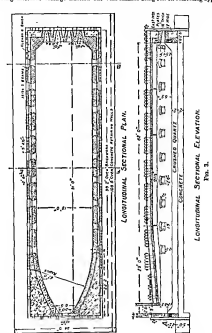
The oxide of iron thus formed, together with that already in the roasted charge, combines with the silica in the roasted concentrates, and in the converter slags added to the charge for copper recovery, together with silica added to the charge to provide the correct amount necessary for slag formation. The copper matte containing all the copper present as sulphide together with any excess ferrous sulphide forms a layer on the bottom of the furnace, and the slag, being of lower specific gravity, floats on the top. The slag is skimmed from the top of the charge and the matte removed from the tap hole as required.

Furnaces 147 ft. long by 23 ft. wide have been used in Anaconda, and with five coal-dust moisture is charged direct to the blast furnace | burners per furnace, 760 tons of charge have been

100 tons of coal dust,

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smelted in 24 hours with a consumption of firms the view that heat transfer in a reverberatory furnace is mainly by radiation. The In pulversed coal or oil firing, the fuel is shape of the furnace has undergone a gradual burnt to give as short a flame as possible and the result is a high temperature peak in the furnace. The resulting increased tonnage ameted con-traction in the flam of the f



a balanced draught. The reactions taking matte autable for recovery of copper and a place in reverberatory furnaces, especially sale of easy furbility and as low in copper as those between iron sulphide, sulphur dravite, possible.

and copper oxides, have been studied by A. (2. TREATHENT OF THE MATTE FOR THE Flat Holdershif (Ind. Eng. Chem. 1930. 22, 256).

Tho object to be attained in this operation in the reverberatory furnace is the same as in euproous sulphile Cuff, with varying proportions the blast furnace, viz. the production of a old ferrous sulphile, FeS, and the further treat-

ment of this matte involves the elimination of the sulphur and iron, leaving the copper in the metallic state. This operation was formerly carried out in two stages in two or more separate furnaces; the first stage consisted in removing the FeS and obtaining fairly pure Cu,S or white metal and involved the partial roasting of the matte leaving only sufficient sulphur to combine with the copper, and the remelting of the whole with silica to form a slag with the whole of the iron present. The second stage consisted in partially roasting the white metal until a certain amount of copper oxide was formed and remelting the whole together, causing the following reactions to take place:

 $\begin{array}{l} \mathtt{Cu_2S+2Cu_2O=6Cu+SO_2} \\ \mathtt{Cu_2S+2CuO} = \mathtt{4Cu+SO_2} \end{array}$

The first reaction is the most important as representing the change which takes place in

processes were mainly carried out in reverberatory furnaces.

The reactions indicated above are now practically always brought about by the "converter" or "Bessemer" process.

The converter consists of a vessel formed of steel plate with a lining of basic material, magnesite bricks, or burnt magnesite. Formerly, a silicious lining was used which took part in the reactions by supplying the silica for the formation of the slag with the ferrous oxide formed. This method necessitated the frequent relining of the converters, which became expensive, so that basic-lined converters are now practically always used, the necessary silica being supplied for each charge. The form of the vessel is more or less cylindrical with the axis of the cylinder placed either horizontally or vertically. In the horizontal form the converter is somewhat barrel-shaped, and a horithe largest proportion of the material. These zontal airbox is attached to one side of the

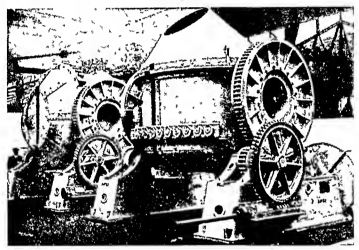


Fig. 4.

vessel for the supply of air to the charge. In the | converter having been tilted into a suitable vertical form (Fig. 4) a shallow circular airbox is attached on the outside at the bottom of the vessel. Air at a suitable pressure is supplied to the airboxes as required and the supply is controlled by suitable valves. The vessel is supported on trunnions or friction rollers and is capable of being rotated by electric or hydraulic power. The vessel is provided with a suitable aperture serving the purpose of a charging opening through which the matte to be treated is poured in a molten condition, also of an outlet through which the products of the operation are poured, as well as being an escape for the gases produced in the process. Charging of the converter with matte is effected by means of a launder direct from the furnace or settler or by tapping these first into a ladle handled by an overhead cranc which conveys it to the con-

In the case of a basic-lined converter, after drying and heating up the lining, a charge of

position for receiving it. The blast is then turned on and the converter tilted over so that the tuyères are brought beneath the surface of the matte and the air, being forced through the molten sulphides, causes a rapid oxidation of both constituents of the ferrous sulphide. The sulphur is oxidised to sulphur dioxide and the iron to ferrous oxide, which is immediately converted into slag by combination with the silica present, forming ferrous silicate. As the oxidation proceeds, the temperature of the charge increases and the operation is continued until practically the whole of the iron sulphide has been oxidised and the contents of the converter consist of cuprous sulphide and slag. The converter is now turned over and the slag run off into ladles and returned for re-treatment in matte furnaces for the recovery of the copper it contains. After pouring off the slag, the converter is again turned into the blowing position when the second stage of the oxidation molten matte is poured in together with the is commenced. The passage of the air through necessary silica to flux the oxide of iron, the the molten cuprous sulphide brings about the

oxidation of a portion of thet material to volatilisation or by passing as oxides into the cuprous oxide which immediately reacts with copper silicate elag formed. The oxidation of unaltered cuprous sulphade producing an the impurities is facilitated by the fact that equivalent quantity of metallic copper in metallic copper has the property of dissolving e accordance with the reactions already given, small proportion of cuprous oxide, which com-This action continues until all the cuprous pound is brought into contact with the elements sulphida has been decomposed and the contents to be exided under very favourable conditions of the converter consist of metallic copper, any small quantity of slag on the surface of the copper is skimmed off for retreatment. The metallic copper is then poured by tilting the converter. As soon as the copper has been poured, the converter is ready to receive another charge and the operations are repeated.

Very large basic lined converters are now being used, and the first stage is often carried out by several separate additions of matte For example, 30 to 40 tons of matte may be poured into the converter followed by 3 tons of silicious ore which may itself be cupriferous. The blust is then turned on, for the first few minutes a pressure of only 5 lb. is used, and after the matte end silica have thoroughly intermingled, of the gases given off by the pole. This process the blast pressure is increased to about 12 lb per sq in When the silica has been fluxed by the iron oxide the converter is turned down and the slag poured off A further charge of when it is ladled out into moulds of any required 10 tons of matte and sufficient adicious ore is now added end the blow resumed removal of slag and additions of matte and silica are continued until sufficient whits metel too long, so that the whole of the oxide present or cuprous sulphide has accumulated in the is reduced, the inguts show a rising along the converter to be blown to metal in the usual centre of the top and the metal is known as

For normal converter prectice matte contain ing 40-45% copper is desired, but the conversion of low-grade matte contening 22% Cu, 45%, Fe, 24%, S is described by H. R. Potts (Trans. Inst. Min. Met. 1928-29, 38, 431) The cherge per blow consists of 21 tons matte. 4 tons flux, and 3 tons copper precipitate from another process. The cherge is blown in a magnesite lined converter, a temperature of I,400° being maintained The blister copper obteined weighs about 5 tons per blow and about 22 tons of slag high in magnetic oxide, and containing 3 4% Cu4S and 0 5% Cu4O are formed.

3. REFINING -The method of treatment to be adopted in this operation depends upon whether (a) the copper is practically free from gold and quantities of deleterious elements

If the metallic copper produced by the under the description (a) above, it can be refining furnace.

Under (a) the furnaca employed is of the

of fluidity, The oxidising condition of the bath of molten copper is maintained until the impurities are removed as far as practicable, and the copper contains a maximum quantity of dissolved cuprous oxide In this condition a cast ingot shows a depression on the surface and the product is known as "under-poled" or "dry" copper After removal of the slag the next stage consists in reducing the greater portion of the dissolved cuprous exide by covering the aurface with carbon and depressing poles of green wood below the surface of the metal, when considerable agitation takes place, and the cuprous oxide is subjected to the reducing action is continued until only a small quantity of euprous oxide remains and the copper ettains the physical condition known as " tough pitch," shape or run out into a large ladie and thence This into mould. In this condition the ingots have a fairly flat surface. If the poling be cerried on " over poled "

The largest refinery plant in this country is at Prescot, where copper is refined for British Insulated Cables, Ltd., from blister copper emported from Rosn Antelope Copper Mines, Ltd. The reverboratory furnaces used are 33 ft long, 13 ft wide (hearth size) and of 190 tons capacity, pulverised coel being used as fuel. Waste heat hoilers are placed at the exhaust end and reduce the temperature of the waste gases to 200°. Fuel consumption for refining and steam raising amounts to 14% of the metal cast, as compared with 50% for refining only, in the smeller furnaces (Mining

J. 1938, 201, 452) Under (b) the copper is refined electrolytically. This process depends for its operation upon the fact that if an electric current is caused to silver and from excessive amounts of deleterious pass through a solution of copper sulphate, elements, or (b) the copper contains appreciable the copper in the solution is deposited at the quantities of gold and alver or contains undue cathodo while acid is set free at the anode. If the snods is composed of metallic copper, then the acid set free at its surface innicipately operations described under (1) and (2) comes dissolves an equivalent portion of copper which passes into solution and is, in turn, deposited refined by furnace treatment only; if it comes at the eathode, and the process thus becomes under (b) it must be refined electrolytically continuous so long as the current passes and in order to recover the gold and silver or to there is copper at the anode to be dissolved. obtain it free from tha impurities, and the Assuming that the copper sulphate solution is electrodes of purified copper are then treated in a properly made up and maintained, and the refining furnace. then only copper will be deposited at the cathode. reverberatory type, and the removal of the while the gold and silver and all the impurities impurities is effected by slowly melting the pigs contained in the copper anodes will either of copper under highly oxidising conditions remain as insoluble slime collected at the and maintaining this condition until the bottom of the vessel in which the operation is impurities are oxidised and removed, either by conducted or will pass into the electrolyte.

slime; other elements, such as iron, zine, and nickel, pass wholly into the solution, and others are to he found partly in the slime and partly in the electrolyte. The cathode surface to start with is a very thin sheet of electrodeposited copper, and when sufficient copper has been deposited on its surface it is removed from the bath and is melted down in reverheratory or other furnaces. The copper to be refined is cast into anode plates ahout 36 in. square and 2 to 3 in. thick, and a number of these are hung in a lead-lined wooden tank and are suitably connected to the positive pole of the electric supply. Alternating with these anode plates are hung the cathode sheets connected with the negative pole. There are two methods of connecting up, viz. "in series," in which the anodes and cathodes are connected in pairs, and "in parallel," in which all the cathodes are connected with the positive pole of the circuit and all the anodes with the negative pole. The parallel system is now generally used. There has been a tendency towards more acid electrolytes than formerly, 180-220 g. of free sulphuric acid with 35-43 g. of copper per litre heing quito common practice. The current density used varies from 1.45 to 3 amp./dm.² The influences of temperature and composition of the electrolyte on the resistance and on the anodic and cathodic polarisation have been determined by E. W. Rouse and P. K. Auhel (Trans. Electrochem. Soc. 1927, 52, 189). The results show that there is a decided drop in the eell voltage with rise in temperature and with increase in acidity. The addition of glue raises the cathodic polarisation, 10 mg. per litre raising the total cell voltage by 60%. The electrolyte is caused to circulate and after it has been used for some time, a certain proportion must be removed regularly for purification. A quantity of anode copper remains undissolved, and when the tank is to be cleaned, this is taken out for remelting and casting into new anodes; the solution is run off and the slime is removed. This latter is washed and dried, any large particles of copper are removed by sieving, and the fine slime is then treated for the recovery of any gold and silver it may contain.

Wet Methods.-With certain ores and residues and under certain conditions, wet methods of extracting copper may he more suitable than smelting methods, and extensive plant are in operation in which new methods or modifications of the old methods are used.

The wet extraction of copper consists of leaching the ore with a suitable solvent either in the natural condition or after roasting and in a suitable state of sub-division, in heaps or in vats fitted with false hottoms and precipitating the copper from the solution by various methods. Wet processes are specially suitable for the treatment of low-grade oxidised ore and for oxidised tailings, hut have also heen successfully applied to the treatment of material containing the copper as sulphide.

At Rio Tinto the massive cupriferous pyrites are treated as mined; large heaps of the mineral

Gold, silver, and lead pass entirely into the lair and moisture and the copper sulphate thus formed is washed out and the copper precipitated on metallic iron.

In some cases dilute sulphuric acid leaching with subsequent precipitation of the copper on iron or hy electrolysis is followed hut for this to he successful the copper minerals must be readily soluble in the acid and the gangue minerals must he insoluble. When carbonates of lime and magnesia are present sulphuric acid leaching cannot be used. When sulphuric acid leaching is used followed by electrolytic deposition of the copper the regeneration of the sulphuric acid yields a solvent suitable for further leaching operations.

Dilute solutions of ammonia associated with ammonium salts have heen used as leaching liquors, as oxide of copper is soluble in such solutions forming copper ammonium compounds, and solutions of cupric ammonium compounds are capable of dissolving metallic copper, yielding cuprous ammonium compounds which are oxidised to cupric compounds by air or other oxidising agents. On distillation the copper ammonium solutions yield ammonia which can he used again and copper oxide which is reduced to metallic copper in reverberatory furnaces.

The method has been used at the Bwana M'Kubwa mine in N. Rhodesia, where the crushed ore is heated to 350° in a gas-fired rotary tube furnace, the product being passed to a closed chamber where it meets a current of producer gas which reduces the copper compounds to cuprous oxide and metallic copper. The material thus prepared is leached with cupric ammonium carbonate solution. Copper and ammonia are recovered from the liquors by boiling. Melting down of the copper precipitate thus obtained yields a marketable product containing about 99.88% Cu.

In the Neil process (Eng. and Min. J. 1908, 85, 556) solutions of sulphur dioxide have been used for the extraction of copper from oxidised or roasted sulphide ores. The ground material is treated in revolving barrels with water, and sulphur dioxide is passed through the mixturo via one of the trunnions. The solution thus ohtained is heated with steam, when ahout 65% of the copper in solution is precipitated as cuprocupric sulphite and the remaining copper is precipitated on iron.

The Longmaid and Henderson process has heen largely used for the treatment of the residues from the pyrites hurnt for the manufacture of sulphuric acid. The residue consists of ferric oxide and is known as hurnt pyrites or blue billy, and may contain up to 3% copper. The process consists in mixing this material with ahout 12% salt and re-calcining, during which process the copper is converted into soluble eupric chloride; this compound is ohtained in solution by lixiviation with water and the copper is precipitated on iron. Any silver present in the pyrites is also obtained in the hrine solution and may he recovered hy the Claudet process.

The original Hunt and Douglas process consisted in heating ores containing carhonates are allowed to oxidise slowly under the action of | or oxides of copper and roasting ores containing 350 COPPEB.

sulphide of copper, as for its success it was flood washing, as the intermediate drying necessary for the metal to be present as cupric periods tend to bring the copper salts to the oxide. The ore was then treated with a solution of sodium or calcium chloride together with ferrous sulphate, which yielded ferrous chleride solution by means of which cupric exide was converted into cupric and cuprous chlorides, both of which are soluble in strong solutions of chlorides:

3CuO+2FeCl₂=2CuCl+CuCl₂+Fe₂O₈

The solution thus obtained was passed over metallic iron for the precipitation of the copper and regeneration of ferrous chleride. The difficulties met with in this process were connected with the tendency to the formation of basic copper salts, the difficulty of separating the solution from the residue and the impossibility of recovering any silver present.

An improved process consists in dissolving the copper oxide by leaching with snlphuric acid, adding ferrous or calcium chleride to the solution to convert the copper sulphate into chloride, and then passing sulphur thouse to precipitate the copper as cuprous chloride:

CuCl_+CuSO_+SO_+2H_O =2CuCl+2H,SO,

The cuprous chloride is separated for further treatment, the sulphur dioxide in solution is eliminated by blowing in hot air, and the residual sulphurio scid is available for the treatment of further quantities of ore.

The Canadian Hydrometallurgical Laboratory has applied the ferme chloride leaching process to high grade iron copper sulphide concentrates | 1935). which have been reduced to mono sulphides by heating in a neutral atmosphere to 600-700°. Precipitation of the copper from tha leach liquors is effected by means of electrolytic iron produced in a later stage of the process The ferrous chloride solution thus obtained is purified by means of ferrous sulphide, then electrolysed for production of electrolyte iron and the ferric chloride re-formed is returned to the lesching tanks (Canada Dept. Mines, Ore Invest. 1927, 138).

W. G. Murray (Chem. Eng Min. Rev. 1929, 22, 52) has shown that ferrous sulphate solution in the presence of air dissolves copper fairly readily from malachite ores but has little action on chrysocolla or enprite. In the absence of air no copper is dissolved from malachite, showing that the solvent action is due to the oxidation of ferrous to ferric sulphate and anbacquent hydrolysis of the latter.

The factors which govern the removal of soluble copper in the leaching of copper ore A. J. Sweet (U.S. Bur. Mines, Tech. Paper 453, more rapid removal of the copper than does being due to porosity. Its m.p. is 1083°, and

surface of the particles very quickly, thus facilitating aubsequent removal by washing.

PROPERTIES .- Copper is the only red metal. ats properties are appreciably influenced by the presence of other elements in the metal, and the

following types of copper are those chiefly used in the industries : High Conductivity (" HC") Copper .- This is

copper of high purity usually containing 99 9% or more of copper. It is used for electrical purposes and where high thermal conductivity

is specially important. Best Select (" BS ") Copper .- This is refined copper containing small quantities of various

impurities usually sufficient to prevent it from quite conforming to the conductivity standard for HC copper. It is suitable for a wide

range of applications. Arsenicel Copper,-This is copper containing up to about 0.5% atsente Its conductivity is lower than that of HC copper. For certain reasons dealt with later, arsenical copper in

widely used for engineering and general purposes. The above types of copper usually contain a small quantity of oxygen and are then known as tough-patch HC, BS, or arsenical copper, For certain purposes, as for welding and pipe manufacture, copper free from oxygen is most suitable and there is available "deoxidised" or "oxygen free" copper of the various grades. In general, the mechanical properties of the above types of copper are not greatly different (Copper Data, Copper Devel Assoc. No. 12,

Copper possesses the properties of malleability and ductility to a very high degree, so that it can be rolled into thin sheets and drawn into fine wire. As a result of cold work the metal becomes harder and eventually brittle, but its malleability and ductility are regained by an annealing operation, and after this it is immaterial whether the metal be slowly cooled or quenched in water. For practical purposes a temperature of about 200° is considered the minimum for annealing pure commercial copper though the presence of small quantities of certam elements, such as silver, nickel or arsenic, necessitates raising the temperature con-siderably. Annealing at 200° is very alow in its effects, and in practice it is more general to anneal for a short time at a temperature of 500-600°. Too prolonged a heating or the use of too high a temperature as to be aveided. For pure commercial copper the tensile strength varies between 10 and 11 tons per sq. in. as cast, between 20 and 28 tons after cold working, have been reported upon by J. D. Sullivan and and between 14 and 15 tons on annealing after work. The elongation varies between 25 and 1929), who recommend that leaching should 30% on 2 in in the east state, between 5 be carried out at night, as the colder solutions and 20% after cold working, and between 50 and then penetrate more rapidly into the particles of 60% on annealing. The Brinell hardness ore, Washing, on the other hand, should be number likewise varies between 40 and 45 carried out in the daytime, as the higher in the cast state, between 80 and 100 after temperatures then ruling favour diffusion cold working, and between 45 and 65 after ontwards of the copper asits. M. Gaggenheim annealing. Its spgr. after working and anneal-and J. D. Sullivan (bid. Tech. Paper 472, 1930) ing is 893, but that of commercial copper show that alternate wetting and drying effect usually ranges from 82 to 88, the lower figures

with the formation of cuprous oxide which dissolves in the metal reducing its freezingpoint to 1,065°, at which temperature the copper-cuprous oxide eutectic solidifies containing 3.5% cuprous oxide. The metal boils at 2,325°. Copper crystallises in the cubic system; well-formed octahedral crystals are found in nature and are sometimes formed during electrolysis. The electrical conductivity of copper is very high and is taken as a relative standard of 100; it compares very favourably with other metals in this respect, being second only to silver with a relative conductivity of 106 and is followed in the scale hy gold with a relative conductivity of 72. The electrical resistivity of copper is 1-7241 microhms per cm.³ at 20°. These resistance and conductivity values apply only to copper after it has been worked and annealed; hars as cast may have a conductivity of 95 and upwards, but complex castings may he as low as 80. The electrical resistance of copper, as of all metals, varies with the temperature and this variation is sufficient to cause the conductivity of HC copper to fall to about 75% of its value at 20°, when the temperature is raised to 100°. The conductivity of copper as of all pure metals is largely affected by the addition of other elements; phosphorus, silicon, iron, arsenic or heryllium when present in quantities less than 1% reduce its conductivity from 100 to 30. The presence of small quantities of silver and cadmium has little effect and for the British Non-Ferrous Metals Research that of oxygen in the amounts present in commercial coppers is negligible. The thermal conductivity of copper is also high, this metal as 100 being only second to silver at 108, and is followed by gold at 76. The thermal conductivity of HC copper is 0.92 g. cals. per cm./cm./second/degree C. at 20°. The specific heat at normal temperatures is 0.092 cals. per g. per degree, but with increase of temperature it rises, the value at 200° being 0.098 cals. coefficient of thermal expansion is 16.6×10-6 per degree. Dry air has no action on copper at normal temperatures, but under ordinary atmospheric conditions the metal becomes slowly tarnished with an oxidised film which changes in time to a mixture containing sulphate and carhonate (v. Co-ordination compounds, this vol., p. 336). This mixed salt deposit is normally green and forms the wellknown patina of copper, and it serves as a protection to the underlying copper. When heated in air a superficial film of oxide is formed which remains adherent and serves as a protection against further attack up to a red heat, but at higher temperatures serious scaling results. Sulphur dioxide when dry either in the gaseous or liquid condition has no effect on copper. When the sulphur dioxide is in a moist, dilute state in contact with the metal it forms a protective covering of hasic sulphate. Carhon dioxide has no action on copper in the dry state. In the presence of moisture the action is very slow and the resulting product, as in the case of atmospheric action, forms a protective skin. Copper is rapidly attacked by chlorine, and ammonia also readily attacks the metal. Ordinary water supplies have no action on copper.

when melted in the air it is rapidly oxidised as an oxidised superficial tarnish soon develops which acts as a protection to the underlying metal. Waters which are acid in reaction may prevent the formation of this protective film and slight dissolution of the copper may result. Nitric acid attacks copper violently, very dilute hydrochloric acid has little effect, hut the concentrated and moderately dilute acid attack the metal, the action being more rapid if the acid he hot. The presence of air is also an important factor in promoting the action. Cold sulphuric acid has little action on copper, but the hot concentrated acid rapidly attacks the metal. Acetic acid has little action on copper, especially in the absence of oxygen, and the metal is used largely for stills, rectifiers, piping, receivers etc., for this acid. Lactic, tannic, formic, and other organic acids have little or no action on the metal. Copper is resistant to dilute solutions of the alkalis, soda, and potash, but not to the action of concentrated solutions. It is fairly resistant to the action of sea-water and of solutions of many salts, but should not he used in contact with inorganic salts which are oxidising agents, such as dichromates, persulphates, perchlorates or ferric chloride. Hydrocarhons, such as petroleum, benzene, etc., have no action, neither have the alcohols, esters, and many other organic materials.

The effects of various impurities on the properties of pure copper have been investigated Association at the National Physical Laboratory, and the following is a summary of the results:

Oxygen has a relatively small effect on the properties of pure copper, and is neither seriously deleterious nor remarkably beneficial. The mechanical properties are not much affected by small quantities and a metal containing 0.1% differs very slightly from pure copper. The electrical conductivity does not fall rapidly with additions of oxygen, and values exceeding 100% of the International Standard are obtained in annealed specimens containing less than 0.1%. 'The relatively small effect of oxygen is due to the fact that the solubility of cuprous oxide in copper is extremely low and may he considered as nil (Hanson, Marryat and Ford, J. Inst. Metals, 1923, 30, 197).

Arsenic.—In the absence of other impurities, copper containing up to 1% arsenic is very difficult to cast and it is difficult to obtain sound castings. A slight amount of unsoundness does not appear to have any harmful effect on the working properties of the metal, which is remarkably malleable and ductile, and can he worked hot or cold to almost any desired extent. It has been found possible by continuous cold-rolling to reduce cast ingots 10 in. in diameter containing up to 1% arsenic to thin strip only 0.004 in. thick. Even when the arsenic is over 7% the ingots were capable of withstanding a considerable amount of cold hammering without developing cracks. Mechanical tests show that arsenic has hut a slight hardening effect on copper; as the arsenic content is raised from 0 to 1.04% the tensile strength of the annealed material rises only slightly, from 14 to 15 tons per sq. in. The

Brinell hardness of annealed arsenical copper much as 10% animony in solid solution, is practically unsflected by the arsense up to Samples containing up to 0.47% can be hot-lyo but the hardness of cold devorted material rolled astafactorily, but those containing of anenne on the electrical conductivity of figure sea be cold-rolled. The presence of the effect of oxygen and very distinctly greater has that off iron. The effect is such that of the same and amount of animony, e.g. up to 0.22%, the containing the same and the containing the same and ameterial. The solubility of areais in solid ness of copper and the fatigue properties are copper in about 7:25%; this solubility changes imposed by the addition of antimony up to try: little with temperature (Hanson and 0.47% hot rolled. With this composition a Marryat, J. Inst. Metals, 1927, 37, 121).

Arsenic plus Ozygen -A series of alloys containing up to 2% arsenic and 0 1% exygen has been examined. The presence of a little naygen greatly improves the casting properties of arsenical copper and no difficulty is experienced in obtaining sound castings with a relatively high density. Oxygen has a harmful effect on the rold working properties of copper which is reduced by the presence of arsenic. Assenical copper is not autable for severe cold-working unless the amenic concentration is at least 10 times that of the oxygen. Copper containing arsenic and oxygen can be rolled hot without difficulty. The mechanical tests of rolled and annealed hars of good quality differ little from those of copper containing the same amount of arsenic without oxygen, provided that the material has not been damaged by over rolling; the tensile strength rises slightly and the elongs. tion falls slightly with increase of areenic content The electrical conductivity is practically identical with that of copper containing the same amount of amenie without oxygen (Hanson and Marryat, shid 1927, 37, 144).

Iron.—Solid copper will dissolve about 4% iron at 1,080°, but the solubility decreases as the temperature falls. Within the limits of solubility, the effect of iron is considerable, especially on the electrical resistance, which increases rapidly as the iron content is raised. When the iron content exceeds about 02% the effect on the resistivity depends largely on working hea between 0.95 and 1.2%, whilst the heat-treatment of the metal. The tensile the limiting content for cold-rolling is 0.79ease. Iron is a decaudiser for copper, but it Small amounts of phosphorus have a pro-

bismuth adversely affect the rolling properties of copper, particularly during borrolling. The shifting amount for hot rolling of bar appears effect occurs in copper containing phosphorus to be less than 0.01% and in all cold rolling when quenched from 690° and tempered at experiments the material fractured when more 400-450°, the maximum effect occurring with experiments the insternal fractured when more 400-450°, the maximum effect occurring with than about 0.05% was present. For severe 0.95% phosphorus; the Binnell hardness cold work hismuth should be excluded from of a quenched rolled strip of this metal

Antimony,-Annealed copper will hold as 43, 41).

asfe range of stress of ±78 tons/in,2 was obtained as compared with ±5 to 6 tons/in. typical of high-quality copper. Antimony lowers the electrical conductivity of copper very considerably but is approximately only half as deleterious in this respect as arsenic (Archbutt and Prytherch, J. Inst. Metals, 1931, 45,

Antimony plus Arsenic,-Within the limits studied 0 05-0 5% each of antimony and arsenic, and under 0 02% oxygen, copper can be hot-rolled without difficulty, so that arsenic assists the hot rolling of copper rontaining antimony. The addition of antimony to arsenical copper improves its tensile atrength both at ordinary temperatures and at 250°. In annealed rod containing 0 49% antimony and 0 059% arsenic, 156 tons per sq in. was obtained at room temperature and 13 tons at 250° With roldrolled atrip containing 0.5% antimony and rolled atrip containing 0.0% animony on 0.53% aresinc a tensile atrength of 37 2 tons/in with 2.3% extension was obtained The fatigue properties of copper containing 0.5% animony and 05% arsenic are similar to those given

above for antimonial copper (idem , ibid. 282),

Phosphorus.—In small quantities phosphorus
produces soundness in, and removes oxides from copper. With 0.04% and more of phosphorus, the densities of cast ropper approach those of rolled bars. The hardness of cast ingots mcreases with increasing phosphorus content. The limiting phosphorus content for hot working hea between 0.95 and 1.2%, whilst strength is raised by 2% icon from 14 5 to 20 tona | 0 95%. Phosphorus, raises the tenseity of per eq. in. Iron has no great embritting effect | relied copper both at ordinary temperature and and copper containing iron can be rolled with at 250° without marked decrease in ductility. ereates difficulties during casting nwing to the innunced effect in raising the fatigue range of formation of films which destroy the con-tinuity of the metal (Hanson and Ford, J. Inst. The softening temperature of cold-rolled copper Ministy of the interest control of the state cold work hismuth should be exemue. It is copied to the purposes 9003% may be regarded as the maximum (Hanson and Furd, Archbutt, and Ford, J. Inst. Metals, 1930,

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656c, 671b

Uses of Copper and its Alloys.—Copper and its alloys have very wide applications in engineering and industries generally, these applications being advantageous owing to the various special properties of the metal. In the first place they are possible owing to the ease of working of the metal; it can be fabricated by cold-working, such as rolling, drawing, pressing, etc., by hot-working, such as pressing, extrusion, forging and stamping, and by machining; it can be joined by welding, brazing, or soldering.

Its high electrical conductivity renders it suitable for cables for transmitting electricity, for windings of electrical machines and apparatus, for electrodes and connections for welding machines and furnaces and for lightning conductors and earthing systems. Its high thermal conductivity makes it suitable for radiators for cooling various liquids, for refrigerator tubes, coils and vessels, for radiators for heating purposes, etc., for locomotive fire boxes, chemical plant, stills, evaporators, etc. Its resistance to rusting and corrosion makes it suitable for various types of chemical plant, brewing plant, roofing, and tubing for water distribution, and some of its alloys for parts of marine machinery, propellers, pumps, etc.

Various alloys in which copper is the chief constituent are largely employed for applications requiring increased strength and other special properties. The most important of these alloys are the brasses or alloys of copper and zinc; other important groups are the bronzes, or alloys of copper and tin, the aluminium bronzes or alloys of copper and aluminium (see ALUMINIUM), and the copper-nickel alloys.

Copper is also alloyed with certain metals to act as a hardener as in the case of standard

silver and standard gold.

Certain elements are added to metallic copper in order to improve its properties, and of these an important class is known as deoxidisers. Tough pitch copper generally contains from 0.025 to 0.08% oxygen, and for making intricate castings, for welding, and for certain other purposes copper as low as possible in oxygen is desirable. The deoxidisers commonly used are phosphorus, silicon, lithium, magnesium, beryllium and calcium; these are added to the molten metal, when they combine with the oxygen and eliminate it, usually by slagging off. To remove all the oxygen it is generally necessary to add a small excess of reagent which remains in the copper and may have an adverse effect on the electrical and thermal conductivity.

Beryllium is not only used as a deoxidiser, but also as a definite alloying element, forming heat-treatable alloys of engineering importance. Copper containing 2.25% Be is known as beryllium bronze; for its heat treatment it is first annealed by soaking at 800°, followed by quenching in water; it is then hardened by a subsequent precipitation treatment by heating for a controlled period at 250-300°. When

DETECTION AND ESTIMATION of copper, see | considerable merit of this material is its high fatigue limit under corrosive conditions.

Copper containing cadmium (approximately 1%) is of importance on account of its combining high strength with conductivity.

Arsenical copper containing 0.3 to 0.5% arsenic is of great importance and most copper products made in Great Britain other than electrical equipment are made from this material. Additions of up to 0.5% arsenic improve the tensile strength of annealed copper by somewhat less than I ton per sq. in., and only slightly increase its hardness value. In the cold-worked condition, however, the tensile strength is increased by 2 tons per sq. in. and the Brinell number is 15 higher. The chief effect of this addition of arsenic is to raise by 100° the temperature at which softening upon annealing first occurs and to slightly increase the strength at elevated temperatures; these two features have been responsible for the adoption of arsenical copper for a large number of applications such as locomotive fire-boxes, etc.

COMPOUNDS OF COPPER.

Oxides .- Six oxides of copper have been reported of which only two are of importance, namely cuprous oxide, Cu2O, and cupric oxide, CuO. Cuprous quadrantoxide, Cu,O, has been considered as being formed by the addition of a solution of copper sulphate to a cooled dilute solution of stannous chloride in caustic potash. It is an olive-green powder which rapidly absorbs oxygen from the air. According to Moser, however, this product is really a mixture of cuprous oxide and metallic copper (Z. anorg. Chem. 1909, 64, 200). Copper trientoxide, Cu.O, is stated to have been obtained as a hard. yellowish mass by heating cupric oxide to over 1,500° (Bailey and Hopkins, J.C.S. 1890, 57, 269). It is probably a solid solution of copper in cuprous oxide. Copper sesquioxide, Cu2O3, has been reported to have been formed by passing chlorine into a strong solution of sodium hydroxide saturated with copper hydroxide (E. Müller, Z. anorg. Chem. 1907, 54, 417). This oxide is not known in the free state. Copper peroxide or dioxide, CuO₂, is not known in the anhydrous form, but in the hydrated form, CuO₂, H₂O, is obtained by allowing finely-divided cupric hydroxide to stand in concentrated hydrogen peroxide at 6° for several days, the mixture being frequently shaken. A yellowish-brown precipitate is formed which, after washing with water, alcohol, and ether, is dried in vacuo (G. Krüss, Ber. 1884, 17, 2593).

Cuprous oxide, copper suboxide or hemioxide. red oxide of copper, Cu₂O, occurs native as cuprite or red copper ore.

It may be prepared (1) by heating fir.ely divided copper in air below a red heat; (2) by gently heating a mixture of 5 parts cuprous chloride and 3 parts sodium carbonate in a covered crucible and separating the oxide by lixiviation; (3) by reduction of an alkaline solution of a copper salt by sugar or certain other organic bodies, as in Barreswil's (Fehling's) heat-treated after cold working, tensile strengths | solution, (4) by heating in a covered crucible a of over 80 tons per sq. in. have been reported. A mixture of 5 parts cupric oxide and 4 parts

copper filings; (5) by the electrolysis of copper; basic natrate and this ignated. E. Müller sulphate solutions under certain conditions; (6) by heating cupric oxide in sulphur dioxide of sulphate: 3CuO+SO₂=Cu₂O+CuSO₃ (Hammick, J.C.S. 1917, 111, 384); (7) free from copper and cupric oxide hy the electrolysis of a slightly alkaline solution of sodinin chloride with copper electrodes (E. Abel and O. Redheh. Z. Elektrochem, 1928, 34, 323)

Cuprous oxide is decomposed by most seids into a cupric salt and metallic copper; bydrochloric acid, however, converts it into cuprous chloride. It fuses at 1,235° and dissolves in molten copper rendering it hrittle or "dry" when present above a certain amount, hut is a most important constituent in many hrands of copper. It is used in the manufacture of ruby glass and for the production of a red glaze on pottery, and, together with the black oxide, forms one of the copper paints used for painting ships' bottoms.

Cuprous oxide has been found to adsorb about 30% by weight of NO, which may be removed at 65-70° by an mert solvent, such as CCi. (Partington, J.C.S. 1924, 125, 72, 663) The oxida has remarkable rectifying and photoelectric properties which are extensively applied in naw types of photoelectric cells (cf. Dubme and Schottky, Naturwiss 1930, 18, 735).

Cuprous hydroxide, hydrated cuprous oxide, Cu,O,zH,O -A yellow form of euprous oxide is obtained by the action of alkali on cuprous chloride or by the reduction of a cupric salt by means of hydroxylamine in presence of alkali. It can also be prepared electrolytically, using an alkalı sulphate as the electrolyte and an anode of pure copper. It may also be formed by the reduction of Barreswil's (Fehling's) solution in the presence of a deficit of tartrate. The yellow precipitate thus formed is not a definite hydrate but is a colloidal form of the oxide with an indefinite amount of water (Groger, Z anorg Chem. 1901, 28, 154, 1902, 31, 326). It quickly changes to an orange or brick red colour and can then be dried unchanged. At 300° it retains only 3%, and at a red heat it loses this water and changes into the red crystalline oxide. The yellow oxide is a powerful reducing agent, especially in ammoniacal solution. This solution, which is colourless, immediately becomes blue on exposure to the atmosphere and thus forms a delicate test for oxygen.

Cupric oxide, copper monoxide, black oxide of copper, CuO, occurs as tenorite or melaconite,

especially at Lake Superior.

Cupric oxide may be prepared by beating the nitrate or carbonate to dull redness or the aulphate to a high temperature. It may also be obtained by heating finely divided copper or and OH' ions, thus: cuprons oxide in sir or oxygen, pure oxygen acts more slowly than air owing to the protective nature of the film of oxide first formed. The oxide used for organic snalyses, (but not for N estimation) may be prepared by moistening copper scale, the mixture of euprous and formation of Cu(OH), 2NH, when only a cupric oxides, with nitric acid and igniting. A small excess of ammonia is present, but of mixture of copper filings with twice its weight Cu(OH), 4NH, with a large excess of ammonia of nitric acid may be exposed to the atmo (Z. anorg. Chem. 1904, 41, 184). A blue aphere until all the copper is converted into solution is obtained by exposing the colourless

prepared cupric oxide by the electrolyais of sodium bydroxide for a long time with a copper cathode either at 12° with stirring or at 60°

without stirring (Z. Elektrochem. 1903, 9, 313). Cuprie oxide is a black powder which agglomerates when heated and fuses at 1,064° forming cuprous oxide. It is slightly hygroscopic especially when in powder form. When heated with organic substances or certain gases it is reduced to metal. The metal produced by reduction in hydrogen always contains that gas, but if the oxide he reduced in formic acid vapour, the metal is free from hydrogen and is suitable for organic analysis (Weyl, Ber. 1882, 15, 1138). It dissolves in scids with the formation of cupric salts. It is soluble to some extent in oils, etc., and it is for this reason that copper cooking vessels require constant cleaning.

Cupric oxide is used in ultimate organic analysis, to make green and blue glass and glazes and to some extent as a pigment. It bas also been used as one electrode of a galvanic cell (Lalande and Chaperon, Compt. rend.

1883, 97, 164).

A colloidal form of the oxlde has been ohtamed by Pasl and Leuze (Ber 1906, 29, 1545), which gives blue-violet solutions, and when dry retains its solubility indefinitely

Cupric hydroxide, CuO, HaO, may be prepared by precipitating a solution of a cupric salt by the addition of alkali and washing rapidly. The hydroxida, which is blue, is hable to turn black during washing owing to the formation of cupric oxide, and when heated with water always turns black. It may be ohtained by passing an electric current through a well stirred solution of potassium nitrate, using a copper plate as anode and one of platinum as cathode (Lorenz, Z. anorg, Chem. 1896, 12, 436)

The hydroxide has been used as a blue pigment by paper stainers but becomes green on exposure to the atmosphere. It corresponds closely with certain highly basic carbonates sold as serdilers

An aqueous solution of sodium hydroxide of ap gr. 1.345-1.370 or of potassium hydroxide of ap gr. 1 453-1:498, will dissolve 0.78 g of euprie hydroxide in 100 c.e., giving a bright hlue solution which yields no precipitate when boiled. If a more dilute alkali solution be used the product is not stable when further diluted and aubsequently boiled (E. Justin-Mueller, Compt. rend, 1918, 167, 779).

Cuprammonum solutions are obtained by digesting freshly precipitated cupric hydroxide in summons solution, and H. M. Dawson and J. McCrae (J.C.S. 1900, 77, 1257) believe that a compound CuO,2NH, or Cu(OH),2NH, is formed which ionises forming Cu2NH,

 $Cu(OH)_1+2NH_2 \rightleftharpoons Cu2NH_3+2OH'$

W. Bonsdorff has also examined the electrical conductivity of solutions of cupric hydroxide in ammonia solution and has confirmed the

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solution of cuprous hydroxide in ammonia to the atmosphere, and cuprammonium solutions are prepared on a large scale by the joint action of oxygen and ammonia on copper filings at a temperature near 0°.

A saturated solution of freshly precipitated and washed cupric hydroxide in concentrated ammonia is known as "Schweizer's reagent" or cuprammonium, it has the property of dissolving cellulose (cotton wool, linen, filter paper, etc.), with the formation of a viscid liquid which has been used for the preparation of "Willesden papers" and for the production of "artificial silk" on a large scale. A. Froment patented the use of strong ammonia saturated with nitrogen which is claimed to dissolve cupric oxide more readily than when free nitrogen is absent (G.P. 139714, 1900), and A. Lecoeur recommends the removal of the nitrites and other crystallisable compounds from the solution by dialysis (F.P. 362986, 1906). The addition of 1-2% of polyhydric alcohols, carbohydrates, gum, etc., to cuprammonium solutions increases their stability and prevents the deposition of cupric hydroxide (Friedrich, B.P. 4104, 1909), and the presence of such substances facilitates the manufacture of the solution. Chattaway has shown (Proc. Roy. Soc. 1908, 80A, 88) that by the reduction of a cuprammonium solution with phenyl hydrazine, copper mirrors can be deposited on glass.

Sulphides.—Cuprous sulphide, Cu₂S, occurs as chalcocite or copper glance. It may be prepared by heating copper to redness in sulphur vapour or by heating a mixture of copper and sulphur. It is also formed as a result of the reaction between metallic copper and certain sulphides, e.g. aluminium, magnesium, cadmium, etc., and E. Schütz has found that copper forms sulphide with 85% of the sulphur in ferrous sulphide when the two are heated together to 1,195° (Metallurgie, 1907, 4, 663). It melts at 1,130° and the sp.gr. of the artificial product is 5·80. It is dimorphous and has a transition

temperature at 91°.

Cupric Sulphide, CuS, occurs as covellite or indigo-copper, sp.gr. 4-60. It may be prepared by heating cuprous sulphide with sulphur at a temperature below the boiling-point of sulphur; by digesting cuprous sulphide with cold strong ntric acid or by precipitating a copper solution with hydrogen sulphide. On the large scale it is obtained as a paste by precipitating a solution of the sulphate with sodium sulphide, and this pasto has been used in calico printing for the development of aniline black.

Several polysulphides of copper have been

reported.

Cuprous Phosphide, Cu₃P, is formed by the interaction of copper and phosphorus, and is present in the alloy known as phosphor-copper, which contains from 9–15% phosphorus. Alloys containing more than this amount of phosphorus have to be prepared in closed vessels owing to the volatility of the phosphorus. They are largely used for the deoxidation of metallic copper and in the manufacture of phosphorbronze. The compound Cu₃P melts at 1,022°, and forms a cutectic with metallic copper which contains 8-2% phosphorus and melts at 707°.

Cupric Phosphide, Cu₃P₂, is formed when phosphine is passed over cupric chloride or when phosphorus is boiled with a cupric salt. When heated in hydrogen it forms cuprous

phosphide.

Copper Arsenides.—Copper and arsenic combine together to form several arsenides, some of which are found to occur as minerals. When copper and arsenic are heated together cupric arsenide, Cu₃As, is formed which freezes at 830°. At 710° there is evidence of the formation of an unstable compound, Cu₅As₂, with 32·1% arsenic (K. Friedrich, Metallurgie, 1908, 5, 529).

Copper Silicide, known as cuprosilicon, Cu₄Si, is obtained when copper and silicon are heated together in an electric furnace (Vigouroux Compt. rend. 1896, 122, 318; Lebeau, ibid. 1906, 142, 154). It can be prepared pure by heating together 17 parts silicon and 9 parts copper in hydrogen and removing the excess silicon by means of sodium carbonate solution. It is silver white with a metallic lustre and is hard and brittle. Its sp.gr. is 7.58 and it is attacked by chlorine and aqua regia. It is used for deoxidising and hardening copper and certain of its alloys (v. Philips, Metallurgie, 1907, 4, 587).

Copper Silicates are formed when exides of copper and silica are fused together. are found in the valious slags obtained during the smelting of copper ores, and especially in those produced in the converters and during the fire-refining of copper. The copper in these slags is readily reduced to metal by carbon in the presence of iron oxide with the formation of the corresponding iron silicate, it is also reduced by heating with metallic iron and is converted into cuprous sulphide when heated with iron sulphide. As hydrated silicates copper occurs two minerals, viz. dioptase or emerald CuSiO3, H2O, copper, and chrysocolla, Cusió, 2H2O.

Copper Carbonates.—The normal carbonate has not been prepared, but double salts of the normal cuprous and cupric carbonates with ammonium and other carbonates have been described.

Some twelve basic carbonates, varying in the ratios of CuO: CO2: H2O have been reported, several of which occur as minerals, the best known of which are malachite, CuCO₃, Cu(OH)₂, and azurite, 2CuCO3,Cu(OH)2; when ground, these minerals form valuable green and blue pigments. The products obtained by the addition of sodium carbonate or other soluble carbonates to solutions of copper sulphate or nitrate are invariably basic, the degree of basicity depending on the conditions of precipitation. These basic carbonates are known as Soda Bordeaux and are largely used as a fungicide. S. U. Pickering (J.C.S. 1909, 95, 1410) states that the ordinary carbonate of commerce approximates to 2CuO,CO₂,H₂O, the formula for malachite, which contains 71·94% CuO, but H. B. Dunni cliff and S. Lal (J.C.S. 1918, 113, 718) found on examination of thirteen samples of commercial carbonate that the amount of CuO varied from 66-16 to 78-60%, and considered product generally approximated that the

of a solution of copper sulphate with a mixture used as a method of preparing pure acctylenc. of sodium carbonate and sodium bicarbonate they obtained a product of fairly constant composition corresponding to

2CuCO,5Cu(OH)...

Many methods have been used for the preparation of the hasic carbonates of commerce, auch as the conversion of scrap copper into basic chlorido by the action of sodinm chloride and sulphuric acid, and treatment of a mixture of this basic chloride and copper sulphate with soda solution, or by agitation of a solution of copper nitrate with chalk, the pasty precipitate obtained is washed and mixed with 8-10% of freshly hurnt lime with the production of a fine blue pigment (serditer), the precipitate without the addition of lime is green.

Salts,-Two series of copper salts are known, cuprous and cupric. The former are mostly insoluble in water and are of little commercial importance They act as powerful reducing agents and readily pass into the copric con dition. Cupric salta are mostly soluble and are of

wide application.

Cuprous Salts .- Cuprous chloride, CuCl. may be prepared by boiling cupric chloride with bydrochloric acid and copper turnings. On the addition of water it is precipitated so white crystalo Cuprous chlorido melts at about 420° into a yellow transparent mass. The dry crystals become yellowish on exposure to light, if moist they acquire a dirty violet tinge.

Cuprous chlorido combines with ammonis to form three compounds, 2CuCl,NH,, 2CuCl,3NH,, CuCl,3NH,. Cuprous bromide and iodido form compounda analogous to the last two (Lloyd, J. Phys. Chem. 1908, 12, 398, cf. W. Biltz and W. Stollenwerk, Z. anorg. Chem. 1921, 119, 97).

The hydrochloric acid solution of cuproua chloride, when exposed to air, absorbs oxygen and acquires a brown colour, aubsequently depositing a pale blush-green insoluble copper oxychloride, CuCl₂,3Cu(OH)₂, similar to the mineral atacamite. An identical aubstance, known as Brunswick green, is prepared by boiling a solution of cupper sulphate with a small quantity of bleaching powder solution, or by exposing freely to air copper turninga moistened with hydrochloric acid.

The use of the exychloride in preference to the sulphate for destroying mould on vines, etc., is recommended by Deletrez (JSCI. 1909, 28, 4381, 200-250 g. per 190 litres of water

are employed

When a hydrochloric acid solution of cuprous chloride and potassium chloride, is treated with acetylene, a yellowish precipitate, C,H,[(Cu,Cl,),KCl], is formed. Other compounds containing cuprous exide or potassium chloride can be obtained by varying the conditions (Chavastelon, Compt. rend. 1901, 132, 1489). Acetylene reacts with solutions of cuprous chloride in ammonia to form a blood red precipitate of cuprous acetylide which on dehydration has the formula Cu,C, and is highly explosive (Scheiber, Ber. 1908, 41, 3816) be obtained as a fused liver coloured, anhydrous

to the formula of azurite; and by treatment; Its decomposition by means of acids has been

Cuprous chloride in a hydrochloric acid or ammoniacal solution absorbs carbon mouoxide forming an unstable compound (Jones, Amer. Chem J. 1899, 22, 287) which under certain conditions corresponds to CuCl,CO,2H,O (Berthelot, Ann. Chim. Phys 1901, [vii], 23, 32).

Cuprous Bromide, CuBr. is obtained by the umon of its elements or by boiling a solution of cupric bromide with copper, and forms a

colourless crystalline mass.

Cuprous Indide, Cul, is the only known include of copper, and is found in the mineral marshite (Broken Ball, Australia). It is obtained as a white precipitate by adding potassium iodide to a solution of copper sulphate, in this way helf the sodine is combined in the precipitate and half is liberated as free ioding; in the presence of ferrous sulphate or sulphurous acid, the whole of the iodine is precipitated as enprous iodide. The fused salt has a spigr, of 5 653 at 15° (Spring, Rec trav. chim. 1901, 20, 79), it melts without decomposition at 628° and is practically insoluble in water.

Cuprous Fluoride, CuF, is prepored by the action of hydrogen fluoride on heated cuprous chloride, the action is complete only when the temperature is between 1,100° and 1,200°, It is also formed by the action of hydrogen fluoride on cupric fluoride, first heated to 600°, and finally raised to 1,100-1,200° fluorido forms a red crystollino mass which melta at 908° It io stable in dry air, but ia converted into cupric fluorido in moist air, at the same time acquiring a blue colour

Cuprous Sulphile, Cu, SO, H,O, 19 obtoined

hy passing culphur dioxide into a hot solution of cuprous acetate in acetic acid. It forms double salts with the alkali oulphites. Cupro cupric

oulphite, Cu,SO,,CuSO,,2H,O, is a red microcryotalline powder,

Cuprous Sulphate, Cu,SO, is obtained by tho action of dry dimethyl sulphate on dry, powdered cuprous oxido (Recoura, Compt. rend. 1909, 148, 1195). A compound of cuprous sulphoto and carbon monoxide, Cu,SO, 2CO,H,O, has been obtained by Joannis (ibid. 1897, 125, 948) Perhard, by adding bydrayylamine sulphate to ammoniscal cupric carbonate, has made ammonio cuprous sulphate Cu.SO.,4NH3 (1bid 1903, 136, 504).

Cuprous Cyanide, CuCN, is obtained by adding potassium cyanido to a colution of copper sulphate, a red precipitate of cupric eyanide is first formed and on boiling evanogen gas is given off and the white cuprous cyanide is precipits ted. Cuprous cyanide forms a number of double eyamdes (Grossmann and von der Forst, Z. anorg Chem. 1903, 43, 94).

Cuprous Theoryanate, CuCNS, is obtained as a white precipitate when potassium thiocyanate is added to copper sulphate solution in the presence of a reducing agent such as sulphur dioxide, ferrous sulphate, etc. This reaction is

need in quantitative analysis. Cuprous Nitrate v. Cupric nitrate.

Cupric Salts .- Cupric chloride, CuCla, may

mass by heating copper in excess of chlorine. In solution it may be prepared by dissolving the oxide in hydrochloric acid. It crystallises in grass-green prisms or needles containing 2 mols. of water which become pale-hlue when dried in vacuo. The anhydrous salt is obtained on the gradual addition of concentrated sulphuric acid to a solution of the chloride (Viard, Compt. rend. 1902, 135, 168). This anhydrous salt is very deliquescent and forms the salt containing 2 mols. of water mentioned ahove. It is soluble in alcohol. When cupric chloride is exposed to the air an oxychloride is formed, and when a small quantity of potash is added to an aqueous solution the oxychloride, 2CuO, CuCl2, 4H2O, is formed as a pale blue precipitate, this turns to a hlack anhydrous powder when heated, hut again hecomes green with absorption of 3 mols. of water when moistened. It is an intermediate product in the manufacture of verditer (v. Carhonates). Some eighteen cupric chlorides have been reported.

Cupric Bromide, CuBr₂, is obtained by dissolving cupric oxide in hydrobromic acid and evaporating in vacuo over sulphuric acid. It is dark-coloured, very deliquescent, and when heated decomposes into cuprous hromide and hromine. It has heen used as an intensifier in photography, for which purpose it is made by mixing equal volumes of CuSO₄,5H₂O (230 g.) and KBr (230 g.) each in 1,000 c.c. H₂O.

Cupric Fluoride, CuF₂, has heen prepared by

the action of fluorine on copper powder, and also hy the action of hydrogen fluoride on cupric oxide at 400°. It forms small white crystals which on exposure to moist air take up water, hecome hlue, and have the composition

CuF,2H,0.

Cupric Sulphate, CuSO₄, is known in the anhydrous form, and also hydrated as CuSO₄, H₂O, CuSO₄, 3H₂O, and

CuSO₄,5H₂O,

the last heing known as blue vitriol, which is found in nature as the mineral chalcanthite, the most important deposits occurring at Copaquire, Chile, where the rock has been found to contain over 12% of the hydrated salt (E. Walker, Eng. and Min. J. 1903, 75, 710). The mineral boothite, found in California, is reported to be CuSO₄,7H₂O (Schaller, Amer. J. Sci. 1904, [iv], 17, 191).

Cupric sulphate may be prepared by dissolving metallic copper or preferably the oxide in sulphuric acid. The solution of copper in sulphuric acid is assisted by the presence of nitric acid or sodium nitrate. On a large scale it is made from metallic copper scrap, etc., by heating this material in a reverheratory furnace, throwing in an excess of sulphur, and closing the doors. When combination is complete the doors are opened to admit air and the temperature kept at dull redness to allow the oxidation of the sulphide to sulphate to take place. The mass, while still hot, is discharged into dilute sulphuric acid, allowed to settle, the solution decanted, concentrated and crystallised. The sulphate so produced is of considerable purity. Copper similarly treated.

When much iron is present as in most copper ores, and especially in copper pyrites, it is impossible to separate the copper and iron sulphates by crystallisation, hecause, although copper sulphate crystallises normally, with 5 mols of water, it forms, in presence of ferrous sulphate, crystals isomorphous with that salt and containing 7 mols. of water. By carefully adjusting the temperature at which the sulphide is roasted, the iron sulphate may be converted into oxide, while the copper sulphate remains almost unaffected. The iron may be removed hy heating the solution of the sulphates to 180° under pressure, when ferrous sulphate crystallises out and may he filtered off. Alternatively, the iron may he separated from the copper by hoiling the solution with a little nitric acid, and then removed as oxide by hoiling with copper oxide or by boiling with lead peroxide with subsequent addition of a little harium carbonate.

When the sulphate is required for purposes in which the presence of iron is not injurious, such liquors may he at once crystallised out with the formation of crystals containing both iron and copper; much of the agricultural copper

sulphate is of this class.

Copper containing gold and silver is sometimes treated with sulphuric acid diluted with its own volume of water, the copper heing thus dissolved while the silver and gold are left in the

residue.

Argentiferous copper ores and mattes are sometimes roasted in reverberatory furnaces, the roasted product heing digested with sulphuric acid until little free acid remains. The solution is then decanted from the precipitated lead sulphate, gold, and other insoluble matter, and passed into lead-lined tanks containing plates of copper upon which all the silver and part of the antimony and arsenic present are deposited, while the greater portion of the hismuth is precipitated as a basic sulphate and the iron is reduced to ferrous sulphate. The liquor is then crystallised, the mother liquors heing used for treating a fresh quantity of roasted material. Copper sulphate is also obtained in the refining of silver by precipitating it upon plates of copper from its solution as sulphate.

An electrolytic process for the production of copper sulphate consists in using a solution of sodium sulphate and copper electrodes, a current of carhon dioxide being passed through the liquid. Copper is dissolved from the anode as sulphate, while sodium carhonate is produced at the cathode. These react, regenerating sodium sulphate and precipitating copper carbonate, which is collected and dissolved in sulphuric

acid (Kroupa, J.S.C.I. 1906, 25, 78).

A process for preparing cupric sulphate from scrap copper has heen patented. In this, the scrap is treated with air in the presence of cupric chloride solution, and the oxychloride sludge so obtained is decomposed with concentrated sulphuric acid, forming copper sulphate crystals and eupric ehloride, the latter heing returned to the process (B.P. 323115).

Copper sulphate crystallises in large, transmatte, copper glance, and other sulphide ores are | parent, blue, doubly oblique, rhombic prisms of sp.gr. 2.28, of the formula CuSO₄,5H₂O.

Solubility of copper sulphate in water at | for some hours in a weak solution of the sulphate different temperatures as g. of CuSO, in 100 g. and ere sown within 24 hours. solution:

t g.	. 12	9	15* 16·1	20° 17-4	25° 18 5	40° 22 8
g.	55° 26 9	60° 28 1		80°	90° 38.5	100° 42 4

When heated, it loses 2 mols, of water at 30°, becoming CuSO4,3H2O, 2 more at 100°, becoming CuSO4,H2O, and at 260° loses nearly all its water; at 360°, however, a little water (0.04%) is still retained, but by raising the temperature gradually to 400° anhydrous autphate is obtained (Richards, Proc. Amer. Acad. 1891, 26, 240).

The salt obtained at 260° is white and hygroscopic, and combines with water with considerable evolution of heat; it combines with water contained in alcohol and other organic liquids with the production of a blue colour, and may be used for detecting the presence of water in these substances, although this test is It bas sp gr. 3 606 and not very delicate begins to decompose at 361° (Vanjukoff, J. Russ. Phys. Chem. Soc 1909, 41, 688). The anhydrous sulphate has also been prepared by heating the pentahydrate at 250° in racuo (F. Krafft, Ber, 1907, 40, 4770)

Copper sulphate absorbe hydrogen chloride with evolution of heat, forming cupric chloride, and liberating sulphurio acid. It is, therefore, used to remove this impurity from each gases as chlorine, carbon monoxide, carbon dioxide, etc. A similar reaction occurs in solution

When mixed in solution with molecular proportions of other sulphates, such as those of zine, magnesium, cadmium, iron, cobalt, manganese, etc., copper sulphate produces welldefined double sulphates which may be rhombec, monochino or trickinic, according to the amount

of water of crystallisation,

A number of ammino-cupric sulphates have heen reported, including Cu(NH₃)₄SO₄, and also Cu(NH₃)₄SO₄, the cu(NH₃)₄SO₄ and below the cu(NH₃)₄SO₄, the cut of the c copper and sodiom, potassium, caesium, rubi-dium, and ammonium are also known, euch as. Na,Cu(SO,),2H,O, K, Cs,Cu(SO,),6H,O, Hb, and (NH,),Cu(SO,),6H,O. K,Cu(SO,),6H,O, Rb,Cu(SO,),.6H,O

Ten basic sulphates of copper have been reported in anhydrous or hydrated forms varying in the ratio of CuO to SO₂, euch as 5CuO_2SO₃5H₂O and 4CuO₃SO₂4H₂O (v. Fowles, J.C.S. 1926, I845).

Copper sulphete is the most important of the copper salts as the large number of copper compounds used in commerce are prepared from this salt. It is largely used in cabco printing. in dyeing, directly or indirectly for the preparaelectro-plating solutions and in galvanic cells It is also used as an antiseptic, as a preventative against rot in timber, and in making preparations for protecting plants from various diseases. In agriculture the crude sulphate containing

A 10-20% solution of copper sulphate, preferably mixed with lime, is applied to vinca with great benefit. A small quantity is found in the grapes and should be removed, although the taumin and sulphur introduced before fermentation remove it in most cases. Karsten has stated (JSC.I. 1896, 15, 367) that attacks of illness resembling dysentery have been caused by a wine which contained sufficient copper to give a visible deposit on steel in 12 hours. Its presence was due to spraying the vines with a copper salt. An effective wash for vines is prepared by boiling 25 parts sodium carbonate and 25 parts resin in 100 parts water; 2 litres of this solution is diluted with 10 litres of water and added to a solution of 2 kilos of copper sulphste in 50-80 litres of water, the mixture neutralised with codium carbonate and made up to 100 litres with water (Perraud, Compt rend 1898, 127, 978).

Copper sulphate is used in preparing copper ferricyanide, a solution of which in potassium citrate solution is used as a toning bath for photographs (Ferguson, "Copper Toning,"

J.S C.I , 1900, 19, 465).

Cupric Nitrite, Cu (NO2)2, is prepared by the double decomposition of cupric sulphate and barrum nitrite. It is thus obtained as a bright green solution. It is said to evolve nitrio oxide even in the cold, and it absorbs oxygen from the air, being slowly changed to nitrete. According to Ray (J.C S 1907, 91, 1405), when the solution us evaporated under diminished prossure over sulphuric acid, nitric oxide is evolved and the residue, hluish-green in colour, consists of a mexture of nitrite and nitrate, insoluble in

A number of double nitrites are known; for example, potsassum copper nitrite, KaCu (NO2)s, is obtained by mixing cold saturated solutions of copper sulphate and potassium nitrite, adding methyl alcohol, filtering, and adding ethyl acetate to the clear green filtrate, when long crystals of the salt separate. Rubidium copper nitrate of similar composition may be obtained in like manner (Kurtenscker, Z. anorg. Chem. 1913, 82, 204). Basic copper nitrites,

3Cu(OH),,Cu(NO,), and 2Cu (OH),,Cu (NO,),,

are also known.

between -20° and +26°.

Cupric Nitrate, Cu(NO3)2, mey be prepared by dissolving metallic copper, the oxide, or the carbonate in nitric acid. The solution at temperatores abovo 26° deposits dark-blue crystals containing 3 mols, of water which melt at 1145° and boil at 170° with evolution of nitric acid and the formation of a green basic salt. This hasic salt may also be prepared hy boiling the solution of the nitrate with copper tion of most pigments containing copper, in or copper hydroxide or with a little alkali. Hydrated crystals containing 9 mols. of water are obtained at low temperatures, -20° to -24°, and crystals containing 6 mols, at temperatures

The anhydrous salt may be obtained by the ferrous sulphate is used to prevent "smut" in action of nitric anhydrido on the hydrated seeds. For this purpose the seeds are soaked salt (A. Guntz and M. Martin, Bull. Soc. chim.

cupric nitrate in liquid ammonia in contact with copper foil J. Sloan (J. Amer. Chem. Soc. 1910, 32, 972) obtained crystals of cuprous ammonio nitrate, CuNO₃,2NH₃, and by allowing the ammonia to evaporate slowly, anhydrous cuprous nitrate is obtained (v. F. Ephraim, Ber. 1919, 52, 236).

Copper Acetate v. ACETIC ACID, (Vol. I.,

p. 54 c).

Copper Arsenite v. Arsenic (Vol. I., p. 477d). Cupric Phosphate, Cu₃(PO₄)_{2,3}H₂O, is produced by the solution of the carbonate in dilute phosphoric acid and beating the solution to 70°. Caven and Hill found that by boiling this normal phosphate with water or by washing with water at 100° until of constant composition the basic phosphate, $Cu_3(PO_4)_2$, CuO, H_2O , is produced, which compound occurs in nature as tbe mineral libethenite (J.S.C.I. 1897, 16, 29). C. O. B.

COPPER GLANCE chalcocite or CHALCOSINE.

COPPER PYRITES v. CHALCOPYRITE. COPRA and COPRA OIL v. COCONUT Ott.

COPROLITES. Concretionary mixtures of calcium phospbate and calcium carbonate, derived from the excrements of certain extinct fishes and reptiles. Used as a fertiliser. Formerly much used in the manufacture of superphosphate of lime.

COPROMESOBILIVIOLIN BILE

PIGMENTS.

COPRONIGRIN v. BILE PIGMENTS.

The COPROSMA. bark, and more especially the roots, of C. grandifolia, C. linariifolia, and C. areolata (Fam. Rubiaceæ), which are widely distributed in New Zealand, possess tinctorial property (Aston, New Zealand J. Sci. Tech. 1918, 1, 3), and being related to madder probably contain either alizarin itself or an allied colouring matter. Thus, an alcoholic extract of C. grandifolia is coloured purple by alkali, and becomes yellow on addition of acid. From the acid solution, by means of ether, an orange-yellow crystalline substance can be isolated in small amount.

A. G. P. and E. J. C. COPTISINE. The roots of Coptis japonica, which are used in China and Japan as stoma-

chics, contain besides berberine two other alkaloids, coptisine (I) and worenine. (I) was first isolated by Kitasato (A. 1926, 1160).

The ground roots are extracted with 50% aqueous EtOH, the concentrated extract acidified with sulphuric acid and the precipitated salts extracted with hot EtOH and H₂O; the sulphate of (I) remains undissolved and is thus scparated from the two other alkaloids (Kitasato, Amer. Chem. Abstr. 1927, 21, 2700).

Coptisine is a quaternary base, only slightly soluble in organic solvents, it is best purified by reduction to the racemic tetrahydrocoptisine, m.p. 217-218° (Spāth, 227-228°), needles from EtOH-CHCl3, and reoxidation with iodine. salts of (I) are crystalline and highly insoluble: (Mitchell, Analyst, 1935, 60, 459).

1909, [iv], 5, 1004). By Icaving a solution of the iodide, yellow needles decomposing above 280°; the *chloride*, orange-coloured prisms, does not melt at 300°. By replacing by -OH groups the methylenedioxy groups by means of phloroglucinol, methylating the phenolic -OH groups, and reducing the compound, racemic tetrabydropalmatine was obtained. Kitasato (l.c. 1926) tberefore attributes to (I) the following constitution:

it was synthesised by Spätb and Posega (Ber. 1929, 62, [B], 1029) starting from tetrahydropalmatine and from protopine (l.c.). Before (I) was found in nature, a base identical with the tetrahydro-derivative had been synthesised by Hawortb and Perkin (J.C.S. 1926, 1769). Späth and Julian (Ber. 1931, 64, [B], 1131) isolated d-tetrahydrocoptisine (m.p. 203-204°, [a]_D¹⁵+310°, in CHCl₃) from Austrian corydalis bulbs. Go (Amer. Chem. Abstr. 1931, 25, 518, or Chem. Zentr. 1931, I, 791) isolated the lævocompound (m.p. $201-202^{\circ}$, $[a]_D^{17} -315^{\circ}$, in CHCl₃) from a Corean corydalis. The alkaloid stylopine, isolated from Stylophorum diphyllum (Fam. Papaveraceæ) by Schlotterbeck and Watkins (Ber. 1902, 35, 7), is probably identical with tetrabydrocoptisine.

COPYING-INK or INDELIBLE PEN-CILS. Originally the pencils sold under these names were intended to be used for copying purposes, and for this reason a water-soluble aniline dyestuff formed an ingredient of the "lead," but the writing done with most of tbem will yield only faint copies. The term "indelible" is also a misnomer, since tbe metbyl violet, which most of them contain, can be readily bleached.

In addition to the soluble dye, the pigment also contains graphite or kaolin or botb. Preparations intended for use in pencils of this type were first patented by Petit (B.P. 4090, 1874), and their composition was described by Viedt (Dingl. poly. J. 1875, 216, 96). Lehner ("Ink Manufacture," 1902, p. 125) gives the composi-tion of four typical Bavarian pencil pigments, made by Faber, and the table at head of page 360 shows the variation in typical pencils determined by Mitchell (Analyst, 1917, 42, 1).

The proportion of graphite in the pigment affects the behaviour of the material towards infra-red rays, and enables many of the pigments to be readily distinguished from one For absorption spectra of (I) and that of the another. Pigments in which there is no graphite tetrabydro derivative, see A. 1927, 1095. The are transparent in an infra-red photograph

Origin	Mois- ture	Residue insol. in water or alcohol		Loss on ignition (gra- phite, etc)	Ash (kaolin, etc.).
American " Austrian British German	2 35	76 45	21 20	27-09	4736
	7-48	51 53	40 99	25-48	31-92
	2 30	60 89	36 81	56-84	4 95
	3 75	63 95	32 30	13-90	50 05
	3 56	47 14	49 94	44-14	3 00
	2 72	65 48	31 80	62-08	3 40
	4 53	73 05	22 42	69-17	3 88
	3 72	64 48	31 80	11-86	52 86
	6 33	59 83	33 84	7-61	52 22
	3 83	57 59	38 58	40-91	17 58

Copying ink pencils containing a large pro-portion of alumina tend to be more indebble than those in which the pigment consists only of graphite and dyestuff. In this country only the violet pencils are in demand, for blue and the votes permit are in command not one such a state of the permit are the permit sould pending an insoluble pigment instead of a dye; the composition of the pigment of the permit of typical coloured pending was determined by Mittebell (Analyst, 1922, 47, 385), Geol Serv., Prof. Paper 124, 1922), and a A ningle charming test, in which the graphite selection of these is given in the table.

of a copying ink pencil is left on the coherent sh of the paper, enables the sequence of two ntersecting atrokes made with a copying ink encil of the graphite type to be determined Mrtchell, Analyst, 1925, 50, 178).

C. A. M.

CORACAN t. CEREALS, (Vol. 11, p. 488d) CORAL is the calcareous substance secreted y several different organisms. The best known If these belong to the Coelenterates and include: I) some Hydrozon known as Millepores. ommon on tropical coral reefs, (2) a large number of Actinozoa (Anthozoa), which include he commonest reef building corals and the secrous coral of industry. The term coral is iso loosely applied to (3) some of the l'olyzon known as sea-mats or zoophytes) which have oral like forms and are the chief constituent of the Tertiary deposit known as the "Coralline Crag", (4) some shallow water Foraminifers, very primitive shell bearing animals which form an important percentage of the sand of coral reefs , (5) certain sea-weeds (Nullipores or Calcareous Alga) which have a calcareous

PERCENTAGE COMPOSITION OF CORAL.

	CsO	MgO	SIO,	Loss on ignition
Hydrozoa— Millepora algicornis	5 2 50	0.43	0 23	41 77
Actinozoa—	02 50	043	0 23	4111
Porties astreoides	53 84	0 18	0 02	44 98
Madrepora prolifera	53 48	0.00	0 19	41 50
Favia fragum	53 69	0.18	0 28	41 44
Maeandra labyrinthiformis	53 71	0 26	(-	45 10
Polyzoa-				1
Flustra membranacea	30 81	206	3 00	61 45
Poraminifera—				1 1
Orbitolites marginalis	48 92	4 93	0.03	45 20
Litholhamnium	48 00	478	0 41	45 41

The precious coral of industry is chiefly Corallium rubrum. It is obtained in depths between 90 and 900 ft. in the Mediterranean The richest fishery is on the banks off the Algerian and Tunisian coasts and also off the coast of Sicily. Black coral comes from Jidds in the Red Sea, and white and red (Morro) coral of good quality comes from Japaneso waters All are used in the manufacture of jewellery. The composition of precious coral is given by G. F. Kunz (in D. K. Tressler, "Marine Products of Commerce," 1923) as shown in table opposite.

	Red coral	Black
Calcium earbonate	86 974 6 804	85 801 6 770
Magnesium carbonate . Calcium sulphate	1.271	1.400
Ferrous oxide	1.720	3 070
Water	0.550	0 600

The colour of black coral is said to be due to gravity of precious coral is close to that of calcite, the organic matter, while some workers have varying from 2 6 to 27, but its hardness (31) is said that the colour of the red cers! salve to a little greater. The low hardness facilitates iron. The red colour, however, is destroyed on carsing and turning and, though it does not ignition, so that presumably it is an organic take a fine polish, the beauty of hie makes up compound formed by the polyps. The specific for this and it wears well. The colour of red being the rarest and therefore most prized.

Although the analyses in the tables give the composition of some of the more important corals, it is known that important changes in the composition take place at comparatively carbonate is largely replaced by magnesium carbonate, which may rise as high as 38%. This imperfectly understood phenomenon is known as *dolomitisation*, and has been most closely studied in the borings from the Atoll of Funafuti (J. W. Judd, Phil. Trans. 1904, "The Atoll of Funafuti," pp. 362-389; E. W. Skeats, Amer. J. Sci. 1918, 45, 185). It was found that down to the depth of 637 ft. dolomitisation bad only occurred to a slight extent, the magnesium carbonate never exceeding 16%; but the lower portions of the bore were very rich in magnesium carbonate, containing from 30 to 43%. Different degrees of dolomitisation have been found in other bores in different reefs.

Sandy islands on coral reefs are the home of innumerable sea-birds, and their excreta replace the carbonate in the coral by phosphates. This guano may be, as is the case on the coral reef known as Christmas Island, of commercial importance. The deposit may contain as much

as 38.5% of phosphoric acid.

(For an account of the distribution, structure, and formation of coral reefs, see J. D. Dana, "Corals and Coral Islands"; C. Darwin, "Coral-Reefs"; W. M. Davis, "The Coral Reef Problem.")

A. P. O.

CORAL-ORE v. CINNABAR. CORAMINE (CIBA). Pyridine β-carboxydiethylamide. Used as a cardiac stimulant.

CORDEAU DETONANT or TONATING FUZE. Several varieties of this fuze, which is used for firing a number of explosive charges simultaneously, have been manufactured, but the type most largely used at present consists of a narrow lead tube filled with trinitrotoluene. It is fired by means of an ordinary detonator and safety fuze or by means of an electric detonator.

H. S. CORDIALS AND LIQUEURS. These are spirituous mixtures usually sweetened and flavoured with essences obtained from herbs, spices or other substances, and often strongly coloured. They are manufactured and used to a much greater extent in France and other Continental countries than in the United Kingdom, and although there is no essential difference between them, the term "cordials" is by many restricted to British and American "liqueurs" preparations, and originating elsewbere.

The spirit in the best quality of liqueur is derived from the grape, but grain, molasses or other spirit is frequently employed, especially in cordials. In the case of citrus cordials the spirit used is often prepared by fermentation of the juice of the fruit after the addition of Whatever the origin of the spirit it should be highly rectified, and unless obtained from the basic fruit it should be neutral in

coral varies from crimson to rosc, the latter | finished product. The alcoholic strength of cordials and liqueurs varies greatly, but, on the average, they contain from 30 to 40% of alcohol, approximately equivalent to 50 to

70% of proof spirit.

The principal flavouring ingredients employed sballow depths in coral reefs. The calcium fall into three classes. Those used solely on account of their essential oils include aniseed, bitter almonds, caraway, juniper berries, mint, lemon-peel and orange blossom. Those valued for their bitter principles as well as their essential oils include cinnamon, cloves, curacoa apples (a kind of orange), gentian, ginger, orange-peel, sweet calamus, wormwood and vanilla. The third class of flavouring consists of fruits, such as cherries, pineapples, raspberries and strawberries. The flavourings are prepared for use either by digesting with alcohol and distilling the mixture to obtain an essence, filtering the alcoholic extract to form a tincture, or by expressing and clearing the juice of fruits. Those for citrus cordials are sometimes prepared by expressing the oil from the peel, the juice of the fruit being fermented, after the addition of sugar, to give the necessary spirit. Artificial essences simulating the above flavourings are frequently employed, but their use in beverages is questionable, since it is possible that their physiological action differs from that of the natural flavouring substances which they are intended to imitate.

The sweetening material used may be sugar, honey or glycerol. In the case of the former it should be commercially pure and, before addition to the preparation, is dissolved in water to form a

syrup.

The colouring matters most commonly used are: red, aniline red, brazilwood, cochineal, cudbear, and sandalwood; yellow, saffron and turmeric; blue, indigo; green, chlorophyll; violet, aniline violet; brown, burnt sugar or caramel.

F. G. H. T.

This smokeless powder, in CORDITE. various modifications, is the principal British Service propellant. It is usually in cord-like form, but for special purposes it is made into tubes with single or multiple perforations, and is sometimes used in the form of flakes

made by slicing the cords.

Composition .- Ordinary cordite is essentially a mixture of guncotton and nitroglycerin, gelatinised by means of acetone, together with a small quantity of mineral jelly (soft paraffin), the whole being thoroughly incorporated. The mineral jelly was originally added to prevent fouling of the rifle, but it was found to diminish erosion by acting as a cooling agent, to improve the regularity of the burning, and also to act as a waterproofing agent. The unsaturated hydrocarbons such as olefines and naphthenes present in the mineral jelly, by combining with the products of decomposition of the nitro-bodies, tend to stabilise the cordite. Furthermore, the mineral jelly to a certain extent protects the nitroglycerin and nitrocotton from atmospheric influences.

The original cordite (Cordite, Mark I) was composed of nitroglycerin 58%, guncotton 37%, character, so as not to impair the flavour of the and mineral jelly 5%. The guncotton had a 362 CORDITE.

nitrogen content of 12-8 to 13 1% and from diameter of the cord, from 2 to 15 days. Cordinabout 10-12% was soluble in sther-alcohol. M.D. requires a much longer time for drying owns to its high nitroglycerin content this than the Mark I type, several months being cordite had a high temperature of explosion, increasers for the largest ages, and produced considerable crosson in genss of large calibre. In 1901 a modified form was greater part of the accione, of which about 18% produced, known as Cordite M.D. In this remains in the pressed cordite. Since the variety the percentage of nitroglycerin was amount of arctone present is appreciable,

Manifacture.—The guncotton containing aretone impregnated as from the stoves through about 30% of moisture is dried in a stove by a a series of towers down which a 30% solution current of hot ar until the moisture is reduced of sodium hisulphite is trickling. The accions to about 0 5%. The cooled, dried guncotton is combines with the hisulplute and when the weighed out into rubber lined canvas hags and solution is nearly saturated it is transferred to a then taken to the nitreglyrerm filtering house, still where after the addition of a little codium where the requisite quantity of nitroglycenn is carbonate the liquor is distilled. The acctone measured out through a rubber tube ou to the thus recovered is purified by fractionation. guncotton. The hag is then carried to the mixing house where the contents are mixed by hand on a lead table which is perforated at one obtain uniformity of composition and hallistic end with \(\frac{1}{2} \) in, boles through which the material properties. This is effected in the case of the is rubbed into the hag below. The product at this stage is known as "Cordite Paste" To effect incorporation, part of the acctone is first poured into a kneeding machine of the Werner Pfleider type, the machine elected, and the cordite paste, together with the remainder of the acetone, about 56% of the weight of the guncotton in all, gradually added. The mixture is covered to prevent loss of acctone and the mixing continued for 31 hours.

At the end of this time the charge of mineral ielly is added and the machine run for a further 31 hours. During these operations the acctone gradually dissolves the guncotten and nitroglycerin and thoroughly incorporates them with the mineral felly. The resulting product is known as "Cordite Dough." The stiff dough becomes somewhat heated during the mixing, but the temperature is not allowed to exceed 40°C., and is controlled by the water jacket with which the machine is provided. The dough is then conveyed in bags to the press house, where it is aquirted or pressed, by plungers working in cylindrical moulds, through a die with one or more holes according to the diameter of the cord required A perforated plate supporting a fine wire gauze is placed above the die to retain mechanical impurities. Pressing into tubes is effected by meane of a die with s pin in the centre, but when small sizes are being pressed considerable trouble is experienced through a partial vacuum being formed and eonsequent collapse of the tube. This can be avoided by the use of a hollow pin with com-

munication to the onter air. The smaller aized cords are wound automatically on to reels as they leave the die, the larger sizes are cut into the required lengths

For drying, the reels of cordito and the in stoyes heated by ateam pupes or by a current of air at a temperature of about 40°C. For the the large sizes at 40°C. varies, according to the to the charge.

considerably reduced, the composition being arrangements are made for its recovery. One of nitroglycern 30%, guncotton 65%, and mineral jelly 5%.

(B. P. 25994, 1901) and convists in passing the

thus recovered as purified by fractionation. After drying is complete, different batches of cordito are thoroughly blended in order to emaller erzes by rewinding several reals simultaneously on to a emgle drum. The larger sizes are blended by hand, the sticks from the drying steve traye being placed in boxes and the contents of these mixed by combining a few from each box, the operation being repeated Before hiending, the sticks are carefully examined and any which are hadly distorted or contain specks of foreign matter are discarded

Properties - Cordite varies in colour from light to dark brown The thin cords of the Mark I material are flexible, but the larger sizes break easily when bent Cordite M D is considerably harder and more brittle and is

difficult to cut with a knife. The epecific gravity of Cordite, Mark I, is about 156-1-57, that of Cordite M D being

1.58-1 59. Cordite is water repellent and can be kept under water, salt or fresh, for some time without its explosive properties being affected. Even when fired wet in a gun the ballistics

enly fall off to the extent due to the evaporation of the water. On prolonged cold storage the nitroglycerin exudes slightly, but is gradually re adsorbed when the temperature rises

Cordite does not ignite very readily, but ence ignited burns relatively slowly with a atrong yellowish flame which is practically smokeless. A etick becomes pointed at the end as it burns and the flame can be extinguished hy blowing strongly on it. A considerable hulk of cordite can be burnt in the open without any explosion. It only burns explosively when strongly confined. If ignition of cordite confined in wooden hoxes occurs, the material does not explode, but burns quietly, although the boxes may be hurst open by the escaping gases. Owing to the difficulty in ignition of cordite a larger sizes on trays are placed on open racks stronger cap of different composition is necessary

to iguite the fine cordite in small arm ammunition than was required for black powder, and smallest sizes no stove heating is required in to ensure the ignition of the larger sizes the the aummer; all that is necessary is to keep charge of cordite has to he primed with an the cordite at a temperature of about 15°C. ordinary black powder or guncotton primer for a few days. The time required for drying which passes on the flash of the igniting tube

Cordite is very insensitive to shock and very varied elimatic conditions have shown that the hallistic stability of cordite is of a high order. The hallistic power is very nearly four times that of hlack powder owing to the increased volume of gases liherated per unit weight and the high temperature of the explosion.

Cordite can give a pressure of more than 120 tons per sq. in. as compared with the maximum of 43 tons obtainable with black powder, but in practice the pressure in the gun does not nearly attain this figure owing to the relatively large size of the gun barrel compared with the

volume of the cordite.

The ballistic properties of cordite are more influenced hy temperature than is the case with black powder, the pressure developed and the velocity of the shot heing greater at increased temperatures, but not to the extent of seriously modifying the shooting under Service conditions. Cordite hurns more quiekly under high pressures than low and the smaller the size of cord the more rapid is the comhustion. The pressure due to cordite is more gradually developed in the gun than is the ease with hlack powder and is hetter sustained.

Numerous analyses of the products of explosion of cordite have heen published by Nohle (Proc. Roy. Soc. 1905, 76, 381). The following results were obtained by exploding the powders in a calorimetric bomb, the loading density

being 0.05:

	Cordite, Mark I.	Cordite M.D.
Vol. permanent gas ml. per g	678·0 877·8 27·15 34·35 17·50 0·30 20·70 100·0	781·8 955·4 % 18·15 42·60 23·15 0·35 15·75 100·0
Composition of total gas: Carbon dioxide Carhon monoxide Hydrogen Methane Nitrogen Water	20·97 26·53 13·52 0·23 15·99 22·76.	14·85 34·87 18·95 0·29 12·89 18·15
Pressure tons/sq. in. Heat evolved (water liquid) cal. per kg. Temperature of explosion (ealculated)	2·9 1,272 5,151°C.	2·7 1,036 4,056°C.

The temperatures being calculated with very cannot he exploded by the passage through low values for the specific heats, are consequently it of a rifle hullet. Exhaustive trials under high and have only relative value. The temperature of explosion as determined hy Rohertson was found to he 2,663°C. for Mark I, and 2,374°C. for Cordite M.D.

The larger volume of gaseous products and the lower temperature of explosion explain why the erosive effect of Cordite M.D. is so much less than that of Cordite, Mark I.

The erosive effects of cordite are less than those of hlack powder and of a different character. With the latter, the hore of the gun is pitted hy the corrosive action of the products of comhustion, whilst with the former only the surface near the powder chamber is smoothly washed away by the passage of the hot gases hetween the projectile and the hore before the inertia of the shot is overcome.

Cordite M.D. gives far less flame than Cordite, Mark I, when used in rifles or machine

Analysis of Cordite Mark I and M.D .-Volatile Matter. 5 g. of the ground cordite are heated for 3 hours on a boiling water plate in an aluminium dish covered with a glass cone, both of specified dimensions (see Marshall, J.S.C.I. 1904, 23, 154).

Guncotton.—2 g. of the ground cordite are extracted with ether for 2½ hours in a weighed Soxhlet thimble. The residue is dried at 82°C.

until constant in weight.

Mineral Jelly.—The ether extract is evaporated to dryncss and the nitroglycerin is removed with methyl alcohol. The treatment alcoholic solution is filtered and any mineral jelly which has been transferred to the filter is washed back into the flask by means of ether. The ether is distilled off and the residue dried in the hoiling water oven, any traces of nitroglycerin vapour being removed by a current of air. The drying operation is continued until the mineral jelly is of constant weight.

Nitroglycerin.—This is usually estimated by

Abel Heat Test .- This is the British official heat test which is applied to nitro explosives generally. The principle of the test is to note the time required to produce a standard tint on a potassium iodide stareh paper when the explosive is heated under specified conditions. The explosive is weighed out into test tuhes about $5\frac{1}{4}$ to $5\frac{1}{2}$ inches long having a capacity of 20 to 22 ml. when filled to a height of 5 inches. They are provided with an indiarubher stopper fitted with a glass rod terminating in a platinum wire hook. On the hook is suspended a test paper about 10×20 mm. in area, the upper half of which is moistened with a mixture of equal parts of glyeerol and water. The test tubes are placed to a depth of 3 inches below the cover of a water bath, the temperature of which is carefully regulated at the specified temperature (usually 160-180°F.). The water bath is a spherical copper vessel about 8 inches in diameter fitted with a cover having a number of circular holes (usually six) and fitted on the underside with elips to take the tubes. The hath is fitted with a thermometer, the bulb dipping 21 inches into the bath, and

CORDITE.

The test is complete when the faint brown in curry powder, and as a carminative. line, which appears at the junction of the wet and dry parts of the test paper which is adjusted to be a of an inch above the cover, is equal in tint to the brown line on a standard tint paper.

CORDITE R.D.B.—Owing to the shortage of acetons during the War a further modification known as Cordito R.D.B. was introduced. This variety contained nitrocotton 52%, nitroglycerin 42%, and mineral jelly 6%, the nitrocotton having a comparatively low nitrogen content (12-12 2%) and capable of being gelatinised by means of ether-alcohol

This cordite was designed to give similar ballistics to cordite M.D., and in practice it was Talsif. 1915, 8, 345) gives the following comfound to be in no way inferior to the latter in this position of a sample of coriander fruit :

respect,

In the case of Cordite R D B, the solvents were originally recovered by absorption in sulphnric acid. The Daniel and Bregeat process (B.P. 127309) takes advantage of the property of phenol and its homologues such as cresol of absorbing ether and alcohol, and this method of solvent recovery was employed at various factories during the War with very satisfactory results, about 50% of the total solvent being recovered.

COROYLITE or BARIUM-PARISITE. Carbonate and fluoride of cerium metals and barrum, (CeF)₂Ba(CO₂)₃, containing Ce₂O₂ 23 72% and (La,Ol)₂O₂ 25 67%. It occurs as amall, pule wax-yellow, elub shaped hexagonal crystals in syenite pegmatite at Narsarsuk near Julianehash in south Greenland, spgr. 431. It is isomorphous with, and closely related to, parisite.

L. J. S. CORÉOPSINE POUR CUIR v. Acatoms DYESTURES (Vol. I, p. 133).

CORIANOER .- The dried fruit (seed) of Coriandrum sativum Linn. (Fam. Umbellifetæ). indigenous to Italy, but cultivated throughout Europe, especially in Russia. The plant is an herbaceous annual with an erect and branching stem rising to about 2 ft. in height. It bears whits or pinkish flowers in June and the fruit ropens in August, the fruits being in desimes contrast to all those other umbelliferous plants employed for flavouring purposes in that they are globular with a diameter about twice the size of mustard seed. The plant is abnormal elso in that any part of the fresh plant, when brused, emits a fortid odour, whist the fruit becomes fragrant on drying. The plant is cut and threshed when the fruit is ripe. The fruit is composed of two pale yellowish-brown suh globular mericarps united at their margins, 2 to 4 mm, in diameter, and in cross-section reveals its similarity to other nmbelliferous fruits, in that each mericarp possesses five in a lanabol, formerly known as corundred primary wavy ridges but with four more. The terpenes present are a-pinene, \$\tilde{p}\$ planes, prominent, straight, secondary ridges Vittes, \$p\$ cyamene, depending, terpinene, t or oil ducts, are only present on the commissural and phellandrene. It also contains traces of sides of the ripo fruit, two in each mericarp; borneol, geramol, n decylic aldehyde, and esters there are many others in the unripe fruit, of sectic and decylic acids, but these disappear as it develops. The taste

usually aome form of temperature regulator, largely used for flavouring purposes, especially

Microscopic Appearance - During ripening the outer persuarp is almost entirely discarded. so that stomata are seldom observed, and the chief diagnostic features are the pitted fusiform cells of the lignified aclerenchyma, which are only present on the outer surface of the mericarp, These are five to ten layers deep and cross each other. The vitte are 300 to 400µ in diameter.
The inner mesocarp merges into the endosperm
and consists of thick walled parenchymatous cells, containing alcurone grains, fixed oil, and minute resettes of calcium oxalate.

Chemical Composition .- C. Arragon (Ann.

Water					92
Ash					7 01
Fat					206
Essent	al.	oil			22
Protest	١.				14 2
Fibre					247
Starch	an	d re	duc	oing	
aubs'	tan	ces			21.2

All the figures other than water are on the

dry material.

The principal constituent is the essential oil, which varies from 04 to 10%, the figure obtained by Arragon (supra) is high, and the best method of determination is probably that of Cocking end Middleton (Quart J. Pharm. 1935, 8, 435). The powdered fruit is mixed with brine, distilled, and the vapours passed through the top of a condenser into a graduated tube, in which the oil is collected whilst the condensed water is returned to the distillation flask. An air inlet or outlet is provided by means of a side tube below the condenser but above the graduated tube. These authors found that the yield of oil as increased slightly by powdering the fruits before distillation, but that the powdered drug loses oil on atorage. They found from 0.25 to 0 40% of oil in commercially powdered fruit. For further details of the oil, see Coriander, Essential Oil or.

Standards .- The "British Pharmacoponia" requires that coriander shall contain not more than 2% of foreign organic matter, ash not more than 7%; acid insoluble asli not more

than 1%.

T. McL.

CORIANDER, ESSENTIAL OIL OF. The oil distilled from the ripe fruits of Cornandrum saturum Linn. (Fam Umbellifers).
a native of central Europe and cultivated in Morocco, Thuringia, Russia and India. also grown in Cyprus. Other parts of the plant yield an inferior oil having a bug like odour. Constituents - The chief odorous constituent

Characters .- A colourless or pale yellow oil, is sweet and aromatic, and the fruits are sp gr. 0 879-0 885, optical rotation + 8° to + 13°,

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ref. ind. 1.456 to 1.478. Soluble in 3 volumes of 70% alcohol. Alcohols determined by formylation 60 to 80%. The determination of linalool cannot be made by the acetylation process owing to the decomposition of the alcohol.

This oil is used largely in the manufacture of liqueurs and in the chocolate industry. It is also employed in medicine as a carminative and

flavouring agent.

C. T. B.

CORIANDROL v. Coriander, Essential OIL OF.

CORINDITE is the name given to a refractory and abrasive material which has been claimed as a "new product." In 1914 a French engineer, Noel Lecesne, patented a process of fusing bauxite economically. mixture of 3 parts crude bauxite, as quarried, and 1 part of anthracite is heated in a cupola oven and air blown into the lower portion. The bauxite is reduced, with the formation of aluminium carbide, which in the presence of excess of air is almost entirely oxidised to alumina, the heat of the reaction being such that the mass fuses in successive layers. On cooling, a vesicular mass is left, through which an immense number of small crystals of fused alumina are disseminated (F.P. 471513; J.S.C.I. 1915, 34, 356). There are two varieties of French bauxites, white and red; the former when fused has a melting-point of 1,950°C., the latter resembles emery and is a powerful abrasive. It is stated that Irish bauxite (Larne), white and low in Al2O3, hitherto of little commercial value, gave a product equal to that obtained from French material. The crushed corindite is mixed with finely-ground refractory binders, such as bauxite, kaolinitic clay, etc., moistened and moulded into shapes which have a very low air-shrinkage. The dried bricks are fired at 1,350° to 1,400°C. and up to 1,750°C. show very little variation in volume. The porosity varies with the mechanical composition of the mixtures and ranges from 9 to 12%. Lime, magnesia, etc., are not suitable binders as they lower the refractoriness. Corindite articles also have great resistance to depreciation and have been recommended for rotary cement kilns, heavy oil and gas engines, coke-ovens, etc. (Bigot, Trans. Ceram. Soc. XVIII, 267; La Céramique, 1917, 20, 41; Eng. and Min. J. 1917, 104, 417; J.S.C.I. 1917, 36, 1094).

CORINTH (INDANTHRENE RK GREY K) v. ANTHRAQUINONE DYESTUFFS Vol. I, p. 420a.

CORIOFLAVINE. An acridine dyestuff

v. Vol. I, p. 133c.
CORIPHOSPHINS are acridine dyestuffs

v. Vol. I, p. 132b.

CORK. Cork is derived from the bark of two evergreen species of oak, Quercus Suber Linn., sometimes regarded as a variety of Q. Ilex (holm-oak) rather than as a distinct species, and Q. occidentalis Gay. The former yields the best cork, and grows chiefly in countries bordering the Mediterranean, whilst the latter occurs on the Atlantic seaboard of Portugal, Spain, and France.

and Portugal, which together produce about four-fifths of the world's output. Algeria and Tunis also contribute to the supply, and smaller quantities are obtained in Southern France,

Corsica, Sardinia and Sicily.

The cork tree is from 20 to 60 ft. high, 3 to 4 ft. in diameter, and attains an age of 150 or sometimes even 200 years. It thrives best on sandy or granitic soils. The quality of the bark depends on the nature of the soil, trees on poor soils producing bark of fine texture whilst those on rich soils give a thicker, coarser and spongy bark. The cork is developed in the cortex by the activity of a meristematic layer of

tissue, the phellogen or cork cambium.

Barking the Trees.—This operation requires great care, as on it depend the life of the tree and the regularity of the cork slabs produced. Two cuts are made round the stem-one a little above the ground and the other just beneath the spring of the main branches. Three or four vertical cuts are made and the bark is stripped off by inserting beneath it the wedge-shaped shaft of the implement used to make the incisions. Care must be taken not to injure the phellogen. The first stripping is effected when the tree is 15-20 years old. The product, termed "virgin cork," is rough, uneven, and woody in texture, and is of little value as it is not sufficiently elastic for use as bottle corks. It was formerly used for the production of Spanish black and for ornamental rustic work. but of late years it has been employed in the form of ground cork for insulating purposes and for linoleum manufacture, After 8-10 years a further layer of cork is removed from the trees; this, known as "secondary bark," is of much better quality than the virgin cork but not so fine in texture as the third and later strippings, which are collected at further intervals of 8-10 years.

After the slabs of cork have been taken from the trees, they are stacked and weighted with heavy stones to flatten them. The cork is then immersed for 5 or 6 minutes in a tank of boiling water which causes it to swell and improves its softness and elasticity; the water also removes tannin and other soluble matter. In Spain the cork is sometimes passed through a fire, whereby it is superficially charred, and is supposed to acquire special qualities. The slabs are finally hydraulically compressed and baled

for the market.

Physical Properties and Uses.—The chief characteristics of cork that render it peculiarly suitable for the various uses to which it is applied are its low sp.gr. (average about 0.16), its compressibility, elasticity and imperme-ability to both air and water, and its low thermal conductivity. In addition to its extensive employment in bungs and stoppers for bottles and other vessels containing liquids, it is used in making mats, soles for shoes, linings, artificial limbs and many other articles.

cork is granulated Waste by suitable machinery, and then ground to powder. The finest dust is used in conjunction with linseed oil for the manufacture of linoleum and other floor coverings. The most important use of The chief sources of supply of cork are Spain | cork waste, however, is as insulating material.

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Many methods have been devised for the other products. It is charred by atrong sulmethod of making cork slabs or boards is known as Spanish black. carried out hy placing the ground waste in irou moulds, subjecting it to heavy pressure and heating it in an oven at about 450-500°. This treatment effects the cohesion of the cork particles without the addition of any extraneous material, and the product is termed "baked slah cork."

Chemistry of Cark.-Cork consists essentially of suberm, but contains about 29% of cerm or cork war (m p. 250°). According to Karrer, Peyer, and Zega (Helv. Chim. Acta, 1922, 5, 853) cellulose is present to the extent of 16-30%. The composition has been studied by Zetzsche and his co workers (Helv, Chim Acta, 1927, 10, 346, 1928, 11, 272; 1931, 14, 632), Kügler (Arch Pharm. 1884, 217), Gilson (bid 1890, 690), Hy Ingle (JSCI 1994, 1197), M. von Schmidt (Monatsh. 1994, 25, 277, 302,

1910, 31, 347)

Gilson saponified cork from Q Suber with 3% alcoholie potash, and isolated from the filtrate 44% of fatty acids, consisting of: "phellonic acid," C22H43O3, mp 102°, a monohasic acid giving a violet colour with zino chloride iodine giving a violet colour with zino entorne todates acolution, yield 89, "subernine acid," C₁₁H₁₀O₂, a some fluid sticky substance, yield 36%; and a small amount of "philosomo acid," C₁₂H₁₄O₂, m.p. 62-63". He considered subern to be a mixture of complex esters and possibly condensa tion products of different acids Scurti and Tomassı (Gazzetta, 1916, 46, [n], 159) have found that phellonic acid is identical with a hydroxy. beheme acid and that suberinie acid is either ricinoleia acid or an isomeride of it

Schmidt expressed the view that suberin was partly composed of glycerides, although pre-viously only small quantities of glycerol had heen found among the degredation products of cork. In order to account for the excess of acids above the amount required to form the glycerides, he suggested that they were present as anhydrides or polymerides. He was further of opinion that young cork contains glycerides only, which, under the influence of air, light, and probably also of enzymes, suffer hydrolysis, the glycerol being partly oxidised to carbon doxide and water whilst the fatty acids partly polymerise and partly form subprintes Zetzsche considers that suberm consists of high molecular polymerides of unsaturated in nature, or because, as in the case of copper, acids, mainly hydraxy-acids, and regards they occur "native" in certain strata from Schmidt's hypothesis that they are present which oxygen and salt solutions are largely as anhydrides as unlikely. On hydrolysing excluded. Structurally, corrosion may affect pure cork (prepared by the action of boiling a metal in several different ways; (1) The aqueous sodium sulphite on crude cork) with changes are confined essentially to the surface, alcoholic sodium hydroxide, Zetrsche and on which the corrosion preduct may either Sonderegger found that sodium phellonate accumulate or he removed in solution or other, and the control of the removed in solution or other, and the control of the removed in solution or other, and the control of the removed in the control of the control of the removed in the control of the contro and introduction of the solution salts were photonic.

(2) The changes at the surface are accompanied by more or less uniercypialline penetration; the total said fraction amounting to 28-30% of this "intercrystalline correson" lead to brittle or the said fraction amounting to 18-30% of the said to brittle or the said to be said to b the crude cork.

manufacture of moulded slahs in which use has phuric acid, but is little affected by hydrochlone been made of various hinding materials, such as acid. Chlorine, hromine, and iodine cause cork gluten, casein, tar, pitch, and mixtures of glue to shrink, the two former first bleaching it, with only and resinous substances. Another On ignition it swells and forms a charcoal

> CORNISH STONE v. CHINA STONE. CORN OIL or MAIZE OIL v. CEREALS.

MAIZE CORNUITE #, CHRYSOCOLLA.

CORONILLIN, C23H23O10, [a]D -176°, the glycoside of the seeds of Coronilla scorpioides and C. varia, is hydrolysed to glucose and an aglucone, C17H21O6. It is not attacked by emulsin, although split by one of the enzymes in Aspergullus niger. The glycoside is very toxic and has a moderate cardisc and low diuretic action (Tanret, Compt rend. 1934, 198, 1637).

COROZO. Vegetable avery the seeds of

the Phytelephas macrocarpa CORRÓSION OF METALS, General. -Corrosion is a chemical process which results in the wastage or deterioration of metals. and may thus he distinguished from crosion, which denotes a wearing away by mechanical agencies. Corrosion may occur either as a direct chemical" or as an "electrochemical" attack, but these are phases of the same funds mental process, the essential feature of which is a decrease in the frea energy of the system. Conditions favouring the first type are more often afforded by contact with gases or moisturecontaining atmospheres, whilst immersion in liquids favours the production of electrical currents and hence the accord type of attack; hut this is hy no means a general rule, and the distinction between atmospheric and summersed corresson is usually dictated by other

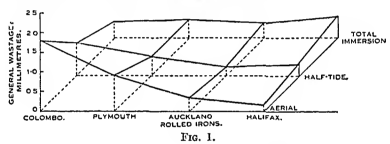
factore to be considered later. The end product of a corrosion process is often a compound which is found in natura as a minerel Thus, copper exposed for many years to an inland atmosphere becomes covered with a coating of basic sulphate which is similar to the mineral brochantite and iron or steel when totally summersed in certain waters and salt solutions, yields, after long periods, hard deposits of hydrated ferric oxide which closely resemble goethite Corrosion is thus the reversion of a metal from a metastable metallic state to a atable ovidised state, metals which occur as such in nature, eg gold, do so because they are corroded only by reagents not commonly found ness or weakness in the underlying metal When treated with strong mitric acid, cork is although the hulk of it may have undergone no converted into auberic acid, exalic acid, and change. (3) The changes are definitely not

confined to the surface, but extend throughout | the mass of the material ("metasomatic change"). In extreme cases a complete change of composition may be effected whilst the article retains exactly its external shape. Examples are provided by the "graphitisation" of cast iron and the so-called "dezincification" of brass, reference to each of which will be made later (see "Corrosion by Sea-water" p. 381 and "Corrosion by Soils," p. 384).

Methods for protection against corrosion should theoretically exclude entirely all substances which promote reversion to the stable condition. These vary with the metal but usually include oxygen, as gas or in solution. water-soluble substances; these are difficult to exclude entirely, and advantage may then be taken of the so-called "self-stifling processes which are described later. A detailed knowledge of the mechanism of corrosion may often provide the key to corrosion protection processes, and an immense amount of work has been done to elucidate this mechanism.

It is convenient to divide the consideration of this subject into two parts, one dealing with metals wholly or partially immersed in liquids,

atmospheres from which liquid water may or may not be deposited; a justification for this procedure is that the factors which control the rates of corrosion in the two sets of conditions are in general different. (By "controlling factor" is understood that factor which, under the particular conditions, has the greatest influence in restricting the rate of corrosion.) For example, it is characteristic of many kinds of atmospheric corrosion that oxygen is present in excess at or close to the metal surface and consequently the rate of corrosion is usually controlled by some factor other than the rate of oxygen supply, e.g. by the humidity, the presence of other gases such as sulphur dioxide, or the nature of the initially formed corrosion product; on the other hand, when a metal is immersed in a salt solution the rate of corrosion is often dependent largely upon the rate at which oxygen reaches the metal surface or its immediate neighbourhood. It is for this reason that differences both in composition of the metal and also in climate give rise, in general, to much greater differences in behaviour on complete exposure to the atmosphere than under conditions of complete immersion in the sea. This and the other with metals exposed to gas is illustrated in Fig. 1, taken from a paper



by Robert Hadfield and S. A. Main Inst. Civil Eng. 1936, 25) in which the authors survey the results of 5-year field tests of the Institution of Civil Engineers. It is apparent from this diagram that maximum differences are exhibited by the aerial tests, maximum agreement by the complete immersion tests, whilst the tests representing "halfconditions occupy an intermediate position.

The surfaces of most industrially important metals exposed to the atmosphere become covered with a film, visible or invisible, which may influence subsequent corrosion processes. This film is frequently an oxide of the exposed metal, and its properties will vary with the metal and the conditions of exposure. It may largely protect the metal from further reaction, e.g. the oxide layer on aluminium which hinders the attack of many salt solutions, and the oxide layer on copper which appreciably slows down atmospheric attack. Of special interest is the resistance of the stainless alloy steels to the corrosive attack of many atmospheres and solutions; this is probably largely due to the protective effect of complex oxide layers formed on their surfaces. On ordinary carbon steels, however, the oxidised layer has little or

(J. 1 exposed to attack by the atmosphere, or by salt solutions or industrial waters. The films formed by exposure to atmospheric conditions may be so thin as to be almost or quite invisible, but heating the metal surface increases the reaction rates and may produce thicker films which show interference colours. Still thicker films formed on steel, for instance, show inherent colours, and may crack and flake away locally during cooling and this may cause a localised electrochemical attack if the metal is exposed to the action of a salt solution. The thick film which remains on the metal acts as the cathode of an electrolytic cell with the bare metal as anode. Then, if there is an ample supply of oxygen to the cathode, and the ions in the electrolyte do not react to form an insoluble, impervious film at either electrode, localised action or pitting may occur at the anode.

Further discussion will now be grouped under two headings, dealing with atmospheric and immersed conditions respectively, to be followed by rather more specific treatment of corrosion in sea-water and in soils. In a survey of this kind it is impossible to give complete references to original work; the references in the text must therefore be regarded only as general steels, however, the oxidised layer has little or guides, and for bibliographical details and more no protective capacity when the metal is freely extended information the reader is referred guides, and for bibliographical details and more

to the following works: "Metallic Corrosion, Passivity and Protection," by U. R. Evans; corresponds to corrosion reta (convenent) London, Edward Aroold & Co., 1937. "Corrosion, Causes and Prevention," by Frank N. the Speller; New York, McGraw-Hill Book Co. 2nd ed., 1935. "Bibliography of Metallic Corrosion," by W. H. J. Veroon; London, Edward Aroold & Co., 1928.

Atmospheric Corresion,

It has been mentioned that etmospheric corrosion comprises reactions of both the metalgas and the metel-liquid type. By the latter is J. Inst. Metals, 1929, 42, 243). The lower range understood any reaction into which water enters, includes the temperatures within which characteristic interests. but not necessarily noe in which the presence of liquid is evident to the eye, Investigations have shown that in eir containing the usual treces of impurities it is not necessary for the meteorological dew-point to be reached for reactions of the accord type to occur; the appropriate conditions are frequently realised well below saturation provided a critical humidity is exceeded, the precise value of which the continuity of which is always mentaned, depends on the metal and on other fectors. The thinner films have been identified by Atmospheric octrosion phenomena may be Derhysheric (Prans. Faradey Soc. 1931, 27, 675), considered, therefore, as felling into three using electron-diffrection methads, as consisting considered, therefore, as felling into three by sing electron-diffraction methods, as consisting groups: (1) The relative below the critical humidity. (2) The relative structure very similar to that of metallic copper, humidity, either continuously or periodeally, and there can be no doubt thet the process is notal is freely exprosed to the open air, and the manual to that vary over the widest possible humidities that vary over the widest possible that hydroxide revour, by direct excitetion, the formation of invalible oxide films, which may sometimes content in the composition of the film. For alimnium and for lead, contation time have a numerotant bearing on the subsections. have an importent beering on the subsequent film; exponential in form in the early stages, course of attack; usually, however, it is the the curves rapidly flatten towards the time axis, mode of breakdown of the film that is of major with which they become nearly perallel in about importance and in this respect the common 10 days, the estimated thickness of film then metals present varied but characteristic being about 10-4 cm. Preston and Bircumshaw features. Both the formetion and breakdown of (Phil. Mag. 1936, 22, 654) have shewn by primary (invisible) oxide films will therefore be electron diffraction that the oxide film on

exposure to air or oxygen. The process may these thin films have within recent years been studied by gravimetric, nptical, photo-electric, and electron diffraction methods.

marked protection against termining (infru) factors that give rise to rusting the oxide film provided that the thickness exceds a critical at ordinary temperatures ceases to thicken, values when appears to be seen that the deals wants and extremely fall, almost immentative lattice of cuprots oried as completed for the after formation. It has, however, been shown whole surface (Vernot, J.C.S. 1925, 2273), (Vernot, Trans. Faraday Soc. 1935, 31, 1668). Rate of navidation increases rapidly with term—that growth of film proceeds continuously, parature, the relationship being given by the although nn ordinary massive metal inter-

ease of metallic exidation. For copper, experimental agreement (linear relationship between log W and 1/T) has been established from 50° to 200° (Veronn, *bid.) and, without change of constants, from 250° to 600° (O. F. Hudson et al., includes the temperatures when when culture teristic interference ("temper") colours are produced, first, accord, and third inders of coloura being discernible. The relationship between exidation and tima throughout both ranges of temperature is given consistently by a quadretic parabola, as would be expected on tha view that thickening of the film proceeds by lattice diffusion of oxygen through an envelope

sluminum at room temperature is emorphous FORMATION OF PRIMARY (INVISIBLE) OXIDE and that crystallisation does not commence until Films -Most metals undergo exidation on the temperature reaches 680°. Zinc is remarkable inasmuch as in ordinary air (free from local be regarded as reversible, se. M+O = MO, pollution but containing the usual traces of the duration depending upon the pressure of impurities) containing its best traces of impurities) containing the best traces of impurities) containing his best traces of impurities) containing the best traces of impurities of containing temperature and for each to the probably associated with however, for have metals excessively low at the breakdown and repair of the primary film-ordinary temperatures and for mallie metals. The man film is granular in structure and does excessively high; this, in fact, constitutes a not show interference colours; eccess of atmo-useful distinction between these classes of aphere to the metal must be assumed to take metals. It is now known that apparently clean place through relatively wide interstices in the surfaces of metals are frequently currend with film instead of by diffusion through economically interesting the properties of timesus envelope (Vernor, Trans, Faraday Soc. I927, 23, 159)

The formation of the invisible oxide film on iron presents a number of distinctive features. The invisible oxide film on copper confers It was formerly thought that in the absence of value which appears to be such that the mut whilst still extremely thin, almost immediately

temperature rises ahove a critical value of very nearly 200°. Below this temperature the film takes on a hrownish appearance which, heyond darkening, undergoes no further change on prolonged exposure. At higher temperatures the first brown or "straw" colour progresses continuously through the familiar sequence of "temper colours," Relationship of oxidation with temperature follows the equation

$d \log W/dT = -Q/RT^2$

ahove and helow the critical temperature, at which there is a change of constants resulting in an increased slope of the log W-1/T curve at the higher temperatures. The oxidation-time curves are quadratic paraholas above 200° but helow this temperature they are modified considerably and correspond approximately with the empirical equation W^{2.5}=Kt. Electron diffraction evidence has confirmed that these changes are associated with a change in the below and aFe₂O₃ ahove 200° (Preston and Bircumshaw, per Vernon, ibid. 1677). See also H. R. Nelson, "The Primary Oxide Film on Iron," J. Chem. Physics, 1937, 5, 252.

Some of the primary oxide films may he removed from the metal by suitable stripping processes. The oxide film produced on aluminium by heating in air was isolated (Seligman and Williams, J. Inst. Metals, 1920, 23, 169) by dissolving away unoxidised metal in hot nitric acid, and that on ordinary unheated aluminium (Withey and Millar, J.S.C.I. 1926, 45, 173T) by volatilising the metal as chloride in a stream of dry hydrogen chloride; a similar method was used (Sutton and Willstrop, J. Inst. Metals, 1927, 38, 259) for isolating the iodide, or by anodic attack of the iron in a solution of sodium chloride; the films obtained were cither fragmentary or complete according to the temperature and time of previous exposure.

Thin oxide films on iron have heen studied spectrophotometrically by Constable (Proc. Roy. Soc. 1927, 115, A, 570; 1928, 117, 376), optically by Tronstad (Trans. Faraday Soc. 1934, 30, 362) using a special technique (reported in a long series of papers) depending on the change in properties of a heam of polarised light after reflection at the metal surface, and electrometrically hy Miley (Carnegie Schol. Mem., Iron and Steel Inst. 1936, 25, 197) who measured the milli-coulombs of electricity required to reduce the film. Valuable contributions have heen made by each of these methods, but the thickness of the film cannot yet be stated with certainty and this hranch of the

sidered hecome of importance in many industrial Faraday Soc., 1938, 34, 867). operations. Pilling and Bedworth (J. Inst. | Relationship with temperature follows in

ference colours do not develop unless the | Metals, 1923, 29, 529) found that the ordinary metals could be grouped in two divisions according to the relation hetween the densities of the metal and the corresponding oxide. Thus, if the oxide occupies a greater volume than the metal from which it is produced, then the film or scale is under a lateral compression which maintains it in a compact and hence protective condition (neglecting possible cracking which may occur under certain conditions), whereas if the oxide occupies a smaller volume its more attenuated condition deprives it of protective value. The first group is represented by most of the heavier metals, typically copper, iron and nickel, the second group by certain of the light metals, typically calcium and magnesium. A further corollary of the continuous nature of the film in the first group is the appearance of interference colours at appropriate thicknesses and also the quadratic relationship hetween film thickness and time. The latter follows hecause the rate of diffusion through the continuous envelope is inversely proportional to the thickness reached, i.e. $dW/dt=k_1/W$, or $W^2=2k_1t+K$, where W represents thickness of film in time t, and K is negligible over a wide range of temperature. On the other hand, films of the second group are characterised by absence of interference colours and by a linear relationship hetween film thickness and time, i.e. $W=k_2t$. Aluminium behaves anomalously for a light metal, hecause its oxide/metal density ratio places it in the first group and its oxide film is in fact extraordinarily protective. On the other hand, it neither exhibits interference colours nor is its oxidation/time curve a quadratic parabola; actually the deviation is in the direction of a cessation of oxidation, the explanation of which is still not altogether clear, although it is ohviously associated with some special property of the alumina film. Preston and Bircumshaw (l.c.) showed that the vitreous film formed at lower temperatures crystallises slowly when heated at 680° to form γAl_2O_3 . In the case of iron the constitution of the scale becomes more complex at higher temperatures. Pfeil (J. Iron and Steel Inst. 1929, 119, 501) has identified three distinct layers, of which the outside layer is lowest and the innermost layer the highest in iron. Pfeil concludes that the oxidation of iron at these temperatures does not proceed by the simple "one-way "diffusion of oxygen through the scale to the metal, but hy a "countercurrent diffusion" in which iron also passes outwards through the scale layer. Pfeil points out that this mechanism does not preclude a parabolic relationship hetween time and oxidation (Portevin, Prétet and Jolivet, infra, found the paraholic law to hold good for pure iron at 825°, 950°, and 1,000°), hut it serves to explain a number of peculiarities in the scaling of iron and iron-rich alloys. The work of Wagner (Z. physikal. Chem. 1933, subject is in active development.

High-Temperature Oxidation.—The properties of oxide films or "scales" formed at and tarnish films (see also L. E. Price, Chem. and Ind. 1937, 56, 769, Hoar and Price, Trans.

temperature films (supra), i.e.

$d \log W/dT = -Q/RT^2$.

(In the case of iron the break in the log W-1/T curve at 200° may be conveniently regarded as "low temperature" and "high temperature" oxidation) The uniformity of this relationship is dependent upon the atmo-ephere remaining unpolluted. For example, minute traces of hydrochloric acid have an extraordinarily marked stimulative effect upon oxidation rate at all temperatures, the mechanism being probably catalytic in character (O. F. Hudson, Herbert, Ball and Bucknall, J. Inst. Metals, 1929, 42, 221). Substances such as carbon dioxide and steam which are neutral at lower temperatures become actively oxidising towards iron and steel at high temperatures, whilst a small amount of sulphur dioxide may profoundly increase the rate of oxidation. The then produces an invisible film of oxide which influence of furoace gases containing these aubstances has been investigated by J. W. Cobb and collahorators, W. H. Hatfield, C Upthe content of reactive sulphur. A similar effect is grove, and other workers (see "Review of produced by preheating in air, when the Oxidation and Scaling of Heated Solid Metals," H.M S O , London, 1935).

The problem of heat treating iron and eteck work in neutral atmospheree (" close annealing ") with the object of avoiding subsequent "pickling" for the removal of scale, has received much attention industrially, and various austable types of furnace are now available The difficulty of this problem arises from the film of reaction product, and the characteristic extremely low pressure of oxygen or other interference colours are in marked contrast to oxidizing gases that is needed for oxidizing to the linear relationship given by zinc, the occur and the difficulty obviously increases con granular nature of the reaction product, and the

aderably with rise of temperature.

Alloys for resisting oxidation at high temperatures most frequently have from nickel or (for the lower limits) copper as the basis metal with additions, most usually, of chromium or aluminium. These metals have a remarkable influence on the oxidation/time relationship, the normal parabolic curve being depressed a continuous tarnish film, whilst the bets towards the time axis. This effect, which (zine rich) constituent shows the discontinuous becomes more marked with mcreasing alloy additions, has been shown for alummium on on zinc (Vernon, Tr. brass by Dunn (J. Inst. Metals, 1931, 46, 42), 886; 1927, 23, 159). and for aluminium and chrominm on fron by Portevin, Pretet and Johvet (Rev. Met. 1934, dependent on the presence of oxygen (Hahn, 31,186). For further information on this subject Z. anorg. Chem. 1917, 99, 118). The loss of the D S.I.R. " Review of Oxidation and Scaling " (supra) should be consulted "TARNISHINO," "Foo

" Focurre," Rusmag (Breakdown of Primary Oxide Films) .-On exposure of metals at ordinary temperatures, copper, silver and mckel alloya have been under sheltered conditions, other phenomena developed by Kenworthy and Waldram (J. besides the formation of oxide films usually Inst Metals, 1934, 55, 247) and W. P. Digby

relative humidities is its susceptibility to tarnish- Chem. (U.S.S.R.), 1935, 6, 601) may play a part ing by extraordinarily minute traces of sulphur in tarmshing processes, since silver and copper compounds, corresponding in their effects to are known to be susceptible to attack by hydrogen sulphide although their precise nature elemental eulphur at ordinary temperatures is nncertain. The reaction is typically of the and the main sulphur containing impurity in metal-gas type; it is greatly facultated by rise air is normally sulphur dionde. Light also of temperature, and is hindered by the presence stimulates appreciably the "fogging" of thin films of moisture. The characteristic mickel (infra).

general the equation already given for low colours are generally believed to be due to the interference of light in a continuous transparent film. Although homogeneous physically, this is composed of a mixture of cuprous exide and cuprous sulphide (extremely similar in lattice structure) with a large preponderance of cuprous oxide. The curve connecting thickness of tarnish film with time is essentially a quadretic parabela; when once started this develops with remarkable regularity and is quita unaffected by wide fluctuations in the degree of atmospheric pollution. In other words, the rate constant is dictated by properties of the initially formed film, which in turn depends on the con ditions of atmospheric pollution which prevailed when the clean specimen was first exposed. An extreme case is presented when at the outset the air is pure or nearly pure-a condition that may be realised during summer months in the absence of local pollution. Exposure for a sufficient tima confera immunity from tarnishing on subsequent exposure to atmospherea containing the usual minimum thickness required for protection (supra) is obtained by an hour's exposure st 75°. The film of oxide serves effectively to keep sulphur atoms out; on the other hand, a thin film of sulphide, hy feculitating entry of both aulphur and oxygen atoms, increases the rate of subsequent tarmshing. The paraholic oxidation/time curve of copper, the continuous granular nature of the reaction product, and the absence of interference colours. The hehaviour of brass as intermediate hetween that of copper and of zine, the oxidation/time curve being compounded of the parabela and the straight line. The microstructure of tarnished 60-40 brass shows that the alpha (copper rich) constituent behaves similarly to copper, giving (granular) structure that characterises the film on zinc (Vernon, Trans. Faraday Soc. 1924, 19,

The tarnishing of eilver by sulphides is reflectivity of eilver and silver alloys in indoor atmospheres is also greatly dependent on the surface condition (Vernon, 1924, shid). Reflectivity methods for following the tarnishing of intervene. The typical cases of copper (with zinc dark present). Go., 1937, 63, 163). The and brass), nickel and iron will be considered. The characteristic feature of copper at low by light (Konstantinova Schlesinger, J. Phys.

attack is controlled mainly hy properties of surface films. Nickel provides an example in which relative humidity hecomes a controlling factor (Vernon, J. Inst. Metals, 1932, 48, 121). In indoor atmospheres the polished surface frequently takes on a creamy-white film, a phenomenon conveniently described "fogging." Here the mechanism is entirely different from that of tarnishing. First, the action depends on the presence of traces of sulphur dioxide as distinct from hydrogen sulphide (supra). But whereas the tarnishing effects of traces of hydrogen sulphide are displayed characteristically at low relative humidities, the corrosive effects of sulphur dioxide come effectively into operation only when the humidity exceeds a critical value in the neighbourhood of 70%. Thus, a polished surface of nickel either retains its hrightness indefinitely or rapidly assumes a creamy film according as the relative humidity remains helow or rises above the critical value. The initial mechanism most probably consists in the catalytic oxidation of atmospheric sulphur dioxide at the metal surface and the reaction of sulphur trioxide with water to give sulphuric acid. In the early stages the film contains free sulphuric acid and may be removed simply by wiping with a cloth; later it passes into a basic sulphate which cannot be removed without ahrasion of the underlying metal. A similar process takes place with copper, but the magnitudes involved are of entirely different orders. With copper a definito concentration of sulphur dioxide has been recognised (approximately 0.9%), which must be exceeded for free sulphuric acid to appear; with nickel, this minimum concentration, although unknown, is excessively low and falls well within the amounts normally Thus, under present in the atmosphere. ordinary conditions, above the critical humidity, the influence of sulphur dioxide is very marked on nickel, but trifling on copper, whilst below the critical humidity the influence of hydrogen sulphide is marked on copper and (within ordinary atmospheric limits) negligible on nickel. Copper-nickel alloys, therefore, may show either one effect or the other, falling off in intensity cither from the copper or the nickel end of the series, according to the relative humidity of the atmosphere.

The rusting of iron presents a number of points of contrast with each of the foregoing processes and constitutes a well-defined third type of attack. It is commonly stated that "iron does not rust ahove the dew-point," hut this is erroneous for most conditions. For example, in an ordinary room atmosphere the relative humidity of which is kept at a low value hy artificial means, an initially clean surface may undergo very considerable rusting. A distinguishing feature is that the attack commences at isolated, sporadically distributed centres; even at a relatively advanced stage the discontinuous nature of the rust can he detected with the aid of a lens. Two further has been allowed to rust in the manner described,

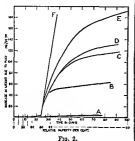
In the processes so far considered, rate of | value (hut still short of saturation) the rate of rusting increases enormously. (2) If the initially clean specimen is screened helind a single thickness of muslin (in form of a " cage ") so as to intercept suspended particles, rusting does not occur and the original hright surface is maintained (Vernon, Trans. Faraday Soc. 1927, 23, 159). It follows from (1) that critical humidity phenomena in the case of iron have a special significance, and from (2) that iron is extraordinarily susceptible to the influence of suspended (disperse) particles of solid or liquid matter in the atmosphere. The screened condition is favourable for the development of the primary invisible oxide film already dis-cussed, and if a specimen has been so exposed for a sufficient length of time, it enjoys a certain degree of immunity from rusting on subsequent normal exposure. After a considerable time-lag as compared with a freshly-cleaned specimen, the primary film hreaks down and rusting proceeds in the usual discontinuous manner.

The influence of both gaseous and solid pollution on the rusting of iron in relation to the relative humidity of the atmosphere has been studied (*ibid.* 1935, 31, 1678) hy experiments in which humidity was increased in steps from zero at the start to a maximum of 99%. The following conclusions are based on these experiments. A similar series of changes occurs either on initially clean specimens in atmospheres containing traces of sulphur dioxide, or on specimens initially "inoculated" with traces of various substances in an otherwise pure atmosphere. In either caso primary and secondary critical humidities are exhibited (Fig. 2). The former, which occurs at 50 to 60% R.H., marks the point at which breakdown of the primary film occurs and attack on the metal commences. The surface hecomes gradually covered with a very fine "rust," heginning at isolated centres, but even prolonged exposure fails to produce severe corrosion; the rust that is formed, however, is in a peculiar physical condition and is potentially extremely active. Thus, at the secondary critical humidity (approximately 80% R.H.) a relatively enormous increase in corrosion occurs, with the production of ordinary red rust. This greatly increased attack is due entirely to properties of the primary rust, for it proceeds just as rapidly if sulphur dioxide (when this is the initiating cause) is completely removed from the system just hefore the critical humidity is reached. The first critical point is associated with optimum atmospheric conditions for the commencement of rusting. The second point must be associated with certain physical properties of the initially formed rust; it is explained by Patterson and Hebhs (Trans. Faraday Soc. 1931, 27, 277) on the basis of Zsigmondy's theory of gel structure. At the lower humidities water is held closely within the capillaries of the gel (the primary rust) and there is a state of mutual compression between rust and water; at the secondary critical humidity the capillaries are filled, the state of compression is released, points must be observed. (1) If, after the iron and water is free to pass from the surface of the rust to the surface of the metal. As a the relative humidity is increased to a high matter of practical interest it may be observed

that the phenomenal increase in attack at this present in urban atmospheres. These several point probably contributes materially to the points are illustrated by the curves in Fig. 2.

The influence of carbon dioxide on the porosity, rusting once occurs beneath it

Particles of solid matter which commonly enter into atmospheric pollution fall, from the corrosion standpoint, into three groups; (a) intrinsically active (dissociating mineral salts) such as ammonium sulphate in the neighbourhood of towns or sodium chloride near the sea; (b) intrinsically neutral but capable of adsorbing active gases from the atmosphere, eg. various forms of amorphous carbon; (c) intrinsically



Curve,	Nature of particles (if any) on specimen. (Where present, uni- formly 04 mg per eq dm)	Nature of addi- tion (if any) to atmosphere (purified alr).
A B C D	None or, charcoal or, quartz Ammonium sulphate None Ouartz	None None
E	Ammonium sulphate Charcoal	Sulphur dioxide

neutral with negligible capacity for adsorption, such as crystalline silica. Only particles of the cold and gaseous pollution are maximal in first two types are important in atmospheric winder and minimal in summer.) The primary corrosion. With the first, the combined presence constituent of atmospheric corrosion products of traces of sulphur dioxide in the atmosphere as in consequence usually the normal metallic and particles at the metal surface produces culphate which, by hydrolysis, may yield either effects at the secondary critical humidity that are hydroxides or basic sulphates, or in some cases almost exactly additive. When, however, the carbonates through contact with carbon dioxide.

particles are capable of adsorbing sulphur Near the seaboard, chlorides or basic chlorides dioxide and so increasing its local concentration may predominate unless a town is near. It is at the metal surface, the effect is more than evident that the attack on the metal may additive; for evemple, charcoal particles have a involve either the removal of a soluble salt or relatively enormous stimulative influence in the the assimilation of an insoluble deposit. To presence of traces of sulphur dioxide. Doubtless denote such processes as these Friend has a similar effect is produced under cervice proposed the terms "crosion" and "surrosion" conditions by carbonaceous particles normally (Fr. eur, on). By crosion is usually understood

atmospheric corrosion of iron has in the past been regarded as of considerable importance. according to the old "acid theory," a trace of carbon doxade permits unlimited attack for the reason that the first-formed product, assumed to be ferrous bicarbonate, hydrolyses to give ferrie hydroxide, with liberation of earbon dioxide, whereby a further equivalent quantity of iron is attacked. Investigation (Trans. Faraday Soc. 1935. 31, 1693; see also Proc. Chem. Eng. Group, S C I, 1937, 19, 19) has, however, abown this view to be untenable. So far from atimulating the corrosion of iron, carbon dioxide, at concentrations in the neighbourhood of and also considerably higher than the normal atmospherio content, has a marked repressive effect at ordinary temperatures This has been abown for a variety of conditions, both "supersaturated" and unsaturated, and including the case where traces of sulphur dioxide (by which the attack is initiated) remain in the system. The effect appears to be directed towards the suppression of the extraordinary change at the secondary critical humidaty. For most conditions the corresion actually observed must depend on the balance between carbon dioxide and sulphur dioxide in the atmosphere. The influence of carbon dioxide on the corresion of sine, particularly under "superseturated" conditions, is in marked contrast. In the absence of sulphur dioxide the effect is repressive, as for iron, although not to such a marked degree; when, however, sulphur dioxide is already present the addition of carbon dioxide stimulates further

OPEN-AIR CORROSION .- Many of the foregoing remarks apply equally to indoor and to open air corrosion, except in so far as the latter is affected by rain. The influence of this factor is not necessarily adverse; it may even be favourable if it removes a corrosion product that is acid or bygroscopic in character, as with nickel, or if rain is able to assist in the development of a product baving protective qualities, as with copper. . On the other hand, the rate of attack may be increased by the breakdown or

corresion.

removal of an otherwise protective coating. In this country rain invariably earnes dissolved sulphates; sulphur dioxide in slight traces is also widely disseminated and may be regarded as a normal atmospheric constituent (Both

as distinct from chemical, processes. Nevertheless, in atmospheric corrosion some distinction is useful because frequently the two processes take place simultaneously, and hence the ultimate change in weight of the specimen or structure is the resultant effect of the two. The endurance of a metal may thus depend on its ability to develop a covering, in inland atmospheres, of basic sulphate, or in marine atmospheres of basic chloride.

The green patina on copper, the corrosion product after exposure (in this country) of some 10 to 20 years, was formerly thought to consist of basic copper carbonate and was attributed to the action of carbon dioxide as present in unpolluted air. Analyses of typical patinas show that in inland districts the product consists essentially of basic copper sulphate, associated with a minor proportion of basic carbonate, the latter being highest in town and least (typically 2.75%) in country samples (Vernon and Whitby, J. Inst. Metals, 1929, 42, 181; 1930, 44, 389; Vernon, ibid. 1933, 52, 93; J.C.S. 1934, 1853). Near the seaboard the sulphate ion in the patina is replaced hy more or less chloride, but when urban and marine conditions coincide basic sulphate greatly predominates over basic chloride. In all cases the patina contains a certain amount of extraneous wind-horne material. Basic copper carbonate is thus accessory either to basic sulphate or basic chloride and is most probably formed by partial hydrolysis of the initiallyformed characteristic salt in the presence of carbon dioxide. Copper is not converted to green basic carhonate, as usually supposed, merely hy exposure to moist air containing carbon dioxide; either the metal must be completely immersed in the aqueous solution or the attack must be initiated in some other way. Small amounts of organic carboxylic acids are particularly active in facilitating the formation of secondary basic carbonate and are probably largely responsible for the relatively high proportion of basic carbonate in urban samples. In districts sufficiently remote from both town and sea (a condition not realised in this country) the green patina does not appear.

The formulæ to be ascribed to the basic salts in the copper patina depend on the period of exposure, maximum basicity being reached only after about 70 years. In the case of basic sulphate and basic chloride 1 molecule of the normal salt is then associated with 3 molecules of copper hydroxide. The formulæ thus correspond with the co-ordination compounds, $[Cu{(OH)_2Cu}_3]SO_4$ and $[Cu{(OH)_2Cu}_3]Cl_2$ each of which is represented in nature by a mineral, brochantito and atacamite respec-tively. Associated basio carbonate has a limiting basicity of only 1:1, as in the mineral malachite, CuČO₃·Cu (OH)₂. Younger products contain higher proportions of normal salt. considerable amounts of which (sulphate or chloride) are leached out in the early stages by the action of rain; it is this process which gives rise to the familiar disfiguration of stonework underlying copper or bronze structures.

the removal of a metal entirely by mechanical, | hefore its final basicity is reached, is insoluble in water, and affords protection to the metal. This accounts for the longevity of copper roofs and suchlike, of which there are many notable examples. There is, however, some evidence that an underlying thin film of cuprous oxide may play a part in the protection. Arsenic is a usual constituent of these old roofs, and this element has been found to reduce the corrosion of copper sheet in the comparatively early stages.

An important directional effect, associated with the influence of prevailing winds, should be noted. Many copper structures have the green patina developed more perfectly on the windward side, particularly if this is directed towards a distant town or towards the sea. A similar principle will obviously apply to other metals in which the corrosion product is not of the protective type, with less desirable effects.

The tendency to build up a protective, insoluble coating on exposure to the open air, very strong in the case of copper when once the early stages have been passed, is usually negligible for ordinary iron and steel. (This statement refers to conditions in this and other industrial countries; but cases are on record where solid and gaseous pollution have been sufficiently low to permit the formation of a protective coating in the course of long exposure.) Under sheltered conditions, depending largely on meteorological and pollution factors, the first-formed rust may actually accentuate subsequent corrosion. Under conditions, however, such that from the outset the air is almost continuously saturated with moisture, an approximately linear relationship with time is followed when once the specimen is covered with rust. Similarly, on prolonged exposuro to the open air, the initial effect observed under less drastic conditions forms only a small part of the total corrosion, and again an approximately linear relationship hetween corrosion and time is followed, the total corrosion being roughly directly proportional to the total amount of pollution that reaches the specimen. The atmospheric conditions prevailing in the very early stages of exposure exert, however, a very great influence in determining the subsequent rate of attack. The importance of this factor in tarnishing has been mentioned. In open-air corrosion it was demonstrated by J. C. Hudson (Trans. Faraday Soc. 1929, 23, 177, 475) for a series of non-ferrous metals which yielded the following mean relative values: summer, 100; autumn, 187; winter, 261; spring, 154. Similar seasonal effects for ferrous materials have been reported by Schramm and Taylerson in the United States (1934, infra), while Daeves in Germany (Naturwiss. 1935, 23, 655) has observed marked differences in country and town atmospheres. G. Schikorr (Z. Elektrochem. 1936, 42, 107) reported on two parallel series of field tests carried out in Berlin and in the Grunewald, 7 km. distant. Fresh specimens of iron were exposed month by month and rate of rusting correlated with metcorological data; hygrometric conditions were very similar, relative On the other hand, the green hasic salt, long humidity in each case being maximal in winter

end minimal in summer. In the purer atmo-| copper reduces corrosion by about 30%, and the apheres the fluctuations in relative humidity joint addition of approximately 0.50 produced comparatively little effect on rusting, copper and chromium by about 40 or 50%, but in the city there was a marked degree of The influence of alloying with larger amounts of parallelism between the monthly corrosion chromium, etc., is discussed under "corrosion. values and the means of the monthly humidity readings. These results ere in harmony with laboratory work which shows that increase of relative humidity, up to saturation, has either a negligible or e profound effect according as gaseous or solid pollution is absent or present.

The influence of climatic conditions has been investigated for marine atmospheres in long period field tests of the Institution of Civil Engineers (specimens of various irons and steels exposed et Halifax, Auckland, Plymouth and Colombo) the results of which have been described by J Newton Friend in a series of reports, the 10 year tests were the last to be described in detail (" Deterioration of Structures trolyte. Exposed to Action of Sea Water," XVth Rep , Exposed to Action of Sea water, Avin Acp, Inst. Cir. Eog. London, I Mis So, 1935 atmosphere, lead, which share with copper a representing up to 15 years' exposure. They have been surreyed generally by Robert resembles copper in developing a protective Madfield and S A. Main (1938, i.e.) Chimatic coating (probably mainly of sulphate) in conditions are also in course of investigation industrial atmospheres and so assuring itself of in a comprehensive series of field tests estah of extreme longevity. In comparison with lished by the Corrosion Committee of the copper, however, it suffers asthetically owing to Iron and Steel Institute and the British Iron the absence of any pleasing patina Contact with and Steel Federation, under the chairmanship new wood (particularly oak), likewise contact of W. H. Hatfield These tests, which are with mortar or cement, should be avoided, or else conducted at fourteen stations distributed in severe corrosion either from organic acids various parts of the world, have been described from the wood, or alkali from the cement may in detailed reports by J. C. Hudson (e.g Vth result. Rep Corr Ctte, Iron and Steel Inst , 1938).

The accompanying data (Table I) are taken from a table given by Hudson (1937, safes).

TABLE I. RATES OF CORROSION OF INCOT IRON AND ZING.

Station	Years'	Rate of sion (1 per y	Ratio.	
	sure	Iron	Zine	
Khartoum	1	42	21	20
Khartoum	5	18	- 8	23
Abisko, N. Sweden	1 1	172	26	66
Aro, Nigeria	1	449	53	85
Basreh, Iraq	1	459	38	12-1
Singapore	1	610	44	139
Apepa, Nigera .	1 1	815	45	181
Redcar .	5	1,590	(534)	(3 0)
Farnborough	5	1,840	89	20 7
Llanwrtyd Wells .	1 1	2,100	118	178
Calshot	1	2,450	133	184
South Africa	1	2,620	174	15-1
Motherwell	1	3,300	185	178
Woolwich	I	3,690	149	248
Woolwich	5	2,000	164	164
Sheffield	1	4,390	579	76
Sheffield University	1	5,040	300	168

In the Fifth Report (supra) Hudson states many phases of the subject were discussed, that the results so far obtained do not indicate a s.e. m 1934 (Proc. Amer. Soc. Testing Materials) marked difference in the corrosion rates of mid and in 1937 (Proc. Chem. Eng. Group, Soc. deem. Ind.).

6 of both resistant alloys" (p. 388). Structural and extraneous factors must be borne in mind. particularly with respect to the incidence of wind and rein; this aspect has been emphasised by Liddierd (Proc. Chem. Eng. Group Soc. Chem. Ind. 1937). Crevices, where water can be retained offer the rest of the structure is dry, are particularly hable to become centres of corrosion, likewise ony place which receives excessive impingement of rain falling from a relatively large estchment area. Contact with non ferrous metals should generally be avoided on account of the danger of enhanced electro-

chemical attack, with rain water as elec-Of other metals commonly exposed to the

Zine is also used successfully for roofing, but much more frequently (e.g., galvanised iron) as a protective covering for iron, over which it has great superiority under most conditions of atmospherie exposure (supra). An acid type of

atmosphere is the least favourable. Aluminium, unalloyed, is resistant to town atmospheres (a good example of its durability is afforded by the Eros statue in Piccadilly Circus, London), but the magnesium aluminium alloy, containing from 2 to 6% magnesium, is advantageous, particularly with respect to the less rapid deterioration of the polished surface Alloye of the duralumin type are habla to severe corresion both in industrial end marine atmospheres, but this may be largely overcome by previous "snodic oxidation" whereby the thickness of the natural oxide film is substantially

mereased Magnesium is noteworthy inasmuch as carbon dioxide enters definitely into the corrosion reaction (Whitby, Trans. Feraday Soc. 1933, 29, 844), for which there is a critical humidity, and carbonate normally predominates over sulphate in the (hydreted) product. Although this

product has no protective qualities, the rate of corrosion, in the absence of elkali chlorides, is not so great as is commonly supposed. The foregoing metals and their olloys ore also discussed under a later heading (p 388)

Reference may be made to the following aymposia on atmospherie corrosion in which

Corrosion in Immersed Conditions.

Although the corrosion of metals immersed in acids, salt solutions, or industrial waters frequently follows an electrochemical course and is proportional to current flow as shown by Evans and Hoar (Proc. Roy. Soc. 1932, 137, A. 355), it is not necessarily so in all conditions. Patten (J. Physical Chem. 1903, 7, 153) has shown that hydrogen chloride dissolved in dry chloroform attacks zine vigorously in spite of the fact that the chloroform has an extremely low conductivity, and experiments have shown that the corresion of steel in "conductivity" water and dilute salt solutions of specially high clectrical resistance must necessarily be largely independent of current flow between attacked and unattacked areas. When the action is electrochemical a large number of factors may influence both the rate and distribution of corrosion. A particularly interesting question is: What are the factors which divide up the metal surface into areas that are anodic (relatively attacked) and cathodic (relatively protected)? Answers that have been given are (1) physical differences in the metal resulting from the mechanical preparation of the surface, (2) discontinuities or points of "weakness" in the air-formed film which overlies the metal, particularly emphasised by U. R. Evans, (3) the presence of different phases within the metal surface, (4) differences of concentration of oxygen, and of ions in solution close to the metal surface, (5) discontinuities such as crystal junctions. It seems probable that only slight differences are required initially to differentiate the metal surface sufficiently to start corrosion. Subsequently, the unequal distribution and the properties of the reaction products may be more important factors in deciding the distribution of anodic and cathodic zones, and this matter is discussed in more detail later. The nature of the corrosion products may also decide the rate of corrosion; in extreme cases this rate may be either reduced to zero by self-stifling at the anodes or allowed to proceed at the maximum rate determined by such other factors as speed of oxygen supply, the concentration of hydrogen ions, etc., detailed in Table II.

TABLE II.

FACTORS WHICH MAY INFLUENCE CORROSION RATES.

Metallic Factors.

- 1. The electrode potential, which is determined by the chemical and physical character of the exposed surface.
- The hydrogen overpotential.¹
 The presence of different phases in the metal surface (especially in acids).

Environmental Factors.

- 1. The temperature.
- 2. Partial pressure of oxygen.
- 3. The rate of supply of oxygen.
- 4. The concentration of H ions.
- 5. The concentration of metal ions.
 - ¹ This term is explained later, p. 376.

6. The concentration and specific nature of other ions present.

7. The electrical conductivity of the solution.
8. The chemical and physical nature and the distribution of corrosion products.

9. Relative movement of liquid and metal.

For a given sample of metal at atmospheric temperatures and pressures the most important factors in approximately stagnant acid solutions are usually the concentration of hydrogen ions, the hydrogen overpotential, the presence of impurities, and the nature of the corrosion product; in many nearly neutral solutions and industrial waters the rate of oxygen supply is also important, and may be so even in dilute acids. In both types of condition the initial corrosive action may commonly be regarded as the passage of metallic ions, at anodic areas, from the crystal lattice into the solution leaving electrons in excess (Me \rightarrow Me"+2 ϵ); simultaneously, an equivalent number of hydrogen ions associate with electrons and leave the solution at cathodic areas on the metal surface $(2H+2\epsilon \rightarrow 2H)$. The hydrogen atoms thus formed may either build up molecules and appear as gas $(2H' \rightarrow H_2)$ or they may react with any oxygen or other equivalent oxidising agent present to form water.

$2H'+2\epsilon+O \rightarrow H_2O$

It will be evident that the depletion of the electrolyte of hydrogen ions will give rise to an excess of hydroxyl ions with consequent alkalinity in the neighbourhood of the cathode. Hoar (Proc. Roy. Soc. 1933, 142, A, 628) prefers to regard the process as a reduction of oxygen at the cathode

$O_0+2H_0O+4\epsilon \rightarrow 4OH'$

but on either view the net result is obviously the same. In weak acids and many salt solutions both hydrogen evolution and oxygen absorption may occur together, gas evolution being predominant in the acids and water formation in salt solutions. These actions can proceed rapidly with relatively base metals such as zinc and iron, and slowly or not at all with semi-noble or noble metals such as copper and gold. Copper yields hydrogen gas only from boiling concentrated hydrochloric acid, but is corroded rapidly in many solutions in the presence of oxygen at ordinary temperatures with formation of water. Gold is attacked only by very powerful oxidising agents such as aqua regia, which evolves chlorine gas, or in the presence of oxygen by solutions in which it forms complex anions. The distinction between the "hydrogen evolution" and "oxygen absorption" types of reaction, notwithstanding many intermediate cases where the processes occur simultaneously, is of much significance and becomes especially important in the consideration of methods of protection.

It is thus useful to consider metals from the point of view of their position in the Electro-potential Series, which is given in Table III. This shows the single electrode potential of each metal when in equilibrium with a solution of a normal concentration of its own ions (i.e.

of normal ionic activity); the potential of the dissolved state. The difference between hydrogen, as exhibited by an electrode of this negative potential and the equilibrium platinum black asturated with hydrogen at an potential is known as the overpotential of pressure of one atmosphere, and plunged in a hydrogen at the surface concerned. It may be solution normal with respect to hydrogen ions, is used as the arbitrary zero point of a "hydrogen scale" in which the potentials of all other metals are arranged. Base metals, i.e. those · below hydrogen in the table, are conventionally regarded as possessing a relatively negative potential, and noble metals a positive one. The base metals can displace hydrogen from many solutions containing its ions, but the noble metals can only do so in special circumatances, e.g. when the concentration of noble metal ions is kept extremely low in the solution by the formation of complex ions containing the metal. The influence of concentration on the single potential may thus be important, and is expressed for any given ignic concentration, C, by the formula:

$$E = E_0 + \frac{RT}{RE} \log_e C = E_0 + \frac{0.058}{R} \log_{18} C$$

at N.T.P., where Eq is the potential at normal ionic concentration, a is the valency of the ion involved, RT and F have their usual meaning, and E is the potential at concentration C.

TABLE III. The Electro potential Series of Metals.

Metal			Ionic state	Potential to volts (on hydrogen scale)	
Gold Platinum .		:	Au Pt probably	+136 +086	
Mercury . Silver Copper . Hydrogen .	:	:	(Hg) ₂ " Ag* Cu" H' (arbitrary	+080 +080 +034 ±000	
Lead Tin Nickel .	:	:	pb" Sn" Ni"	-0 12 -0 14 -0 23	
Cadmium . Iron Chromium Zinc	:	:	Cd·· Fe" Cr·· Zn"	-0 40 -0 44 -0 56 -0 76	
Aluminium Magnesium	:	:	Mg"	-1·33 -1·55	

Arnold & Co , 1937.

The expression " hydrogen overpotential " which has already been used may now be or, perhaps, to a local high concentration of explained. Hydrogen gas is freely evolved from a surface of platinum black whenever the potential is made appreciably more negative ments are those in which the metals are freely potential is made appreciasor more arguered sense are smooth in which the measure of them 00 on the hydrogen scale; from the soluble and no accondary insoluble products are surface of other metals, however, a notably formed; in other solutions and in industrial more negative potential is necressary before waters these conditions may not hold, and if they hydregen is produced either in the gascous or do not no opinion can be gleaned from the

an important factor in determining the rate of corrosion in an acid; for instance, a metal with a high overpotential, such as zinc, will evolve hydrogen and corrode slowly or not st all if chemically pure ; but if it contains impurities of low overpotential embedded in its surface it will corrode more quickly, since hydrogen will be evolved in a continuous stream from the surface of such impurities. These may consist of another metal, alloy, metallic com pound, or some form of carbon. Metals which exhibit low overpotentials include platinum, gold, palladum and iron, while mercury, zinc, lead and tin show the opposits phenomenon; the precise effect of any given metal, however, depends partly upon its surface condition (smoothness, etc).

The cause of overpotential has been tha subject of much work and discussion, and seems to be connected with the sluggishness of one

or both of the reactions .

Hydrogen ion+Electron → hydrogen (atom)

or 2 Hydrogen (atoms) - hydrogen (molecule) Referring to Table III, it will be noticed that the more generally chemically reactive metals have high negative potentials and occur at the bottom of the table, whereas the chemically aluggish metals occur at the top It might be deduced from this that the position of a metal in the potential series would give direct informs tion about its "corrodibility" relative to other metals in the series. The deduction would not necessarily he true, because the rate at which a metal corrodes, s.e. its corrodibility, is partly determined by factors independent of the metal itself, such as composition and mevement of the hound in which it is placed, and the rate of oxygen supply; if, therefore, a metal be placed in a solution other than that in which the determination of its single electrode potential was mada, both its position in the series and its rate of corresion may be altered.

From what has been and above it will have become clear that corrosion rates are largely determined by polarisation at one or both electrodes. Polarisation at ateel cathodes may be due to films deposited from pertain salts in solution (e y. the aulphates and nitrates of zinc and lead) which either shut off the supply of oxygen or other cathodic depolariser, or porson the catalytic activity of the metal surface for the oxidation of hydrogen to water (U. R. Evans, (This table is a condensed form of Table XXIV JS C.I. 1924, 43, 321T; Thornhill and Evans, in "Metallic Corrosion, Passivity, and Pro Fith Corrosion Report to the Iron and Sixel tection," by U. R. Evans. London, Edward Institute, 1933, 381). Polarisation at anotes may be due to the formation of "insoluble" bodies in close contact with the metal surface metal ions at that aurface.

The solutions chosen for the potential measure-

table as to the relative corrosion rates of the To take as concrete examples the three metals iron, magnesium and zinc, the following is the order of their corrosion rates in the liquids mentioned, in presence of oxygen:

Strong KOH. Distilled water. N/10 KCI. Mg Zn Zn Fe Mg Mg

The top metal in each column is the most rapidly corroded and it will be noticed that any one of the metals may occupy any position in the list according to the selection made of the corroding liquid. The general explanation is that the rate of corrosion depends largely on the properties of the corrosion products; thus, in distilled water the hydrated oxide formed affords less protection to iron than it does to magnesium or zinc, but in strong alkali the zinc oxide is dissolved and the metal is readily attacked, while the iron and magnesium oxides are almost insoluble and form films on the metal surface which are highly protective, at any rate for a considerable time. Iron may thus be said to become "passive" in strong alkali as well as in nitric acid and chromate and bichromate solutions of fairly high concentrations. The passivity in these solutions seems to be mainly due to the formation of films of ferric oxide on the surface of the metal; highly protective films are formed only when the precipitation of the metal as oxide occurs at the moment of its entrance into solution, and the essential condition for this is the presence of excess of the precipitating agent at the metal surface. Passivity has been specially studied by Evans and colleagues (J.C.S. 1927, 1020, and subsequent papers) and has been shown to occur with other metals and solutions, when the essential condition is fulfilled of the production of an insoluble product (which is not necessarily an oxide) immediately at the metal surface and in the necessary physical state. Some examples of passivity may be regarded as the result of direct chemical oxidiation, others as caused by complete anodic polarisation. In general, a metal will only remain passive as long as it is immersed in the passivifying solution; if removed to another liquid it will usually corrode because of a lowering of the concentration of the requisite precipitant at the metal surface.

The influence of temperature on corrosion in neutral aqueous solutions depends in general on wbether the system is open or closed. In open systems, as shown by Heyn and Bauer (Mitt. Kgl. Mat. Pruf. 1910, 28, 62) and by Friend (Iron and Steel Inst., Carn. Sch. Mem. 1922, 11, 13), the rate of corrosion of iron increases to a maximum in the neighbourhood of 80°, beyond which the normal stimulative effect of temperature is more than offset by the retarding effect due to expulsion of oxygen. In closed systems, on the other hand, the rate of corrosion increases continuously with rise of temperature (Speller, "Corrosion, Causes and Prevention," 144). Nevertbeless, if entry of oxygen is prevented, this initial rate must fall off rapidly

Thus, the total corrosion can only become serions if oxygen is replenished through any cause. Corrosion at higher temperatures, in strictly closed systems, when oxygen initially present has become exhausted, may still proceed with liberation of hydrogen; the rate at which this reaction occurs, however, is usually so very much lower (with iron and steel and neutral or nearly neutral waters) that it cannot give rise to serious corrosion, although the presence of hydrogen from this cause may frequently be recognised in heating

The influence of light on corrosion, notwithstanding its theoretical interest, does not in general constitute an important factor in corrosion problems. Experimentally it is frequently difficult to separate the effect of light from that of temperature. General observations concerning the influence of light on the corrosion of iron in aqueous solutions have been made by Friend and others (Carnegie Schol. Mem., Iron and Steel Inst. 1922, 11, 116). Reference bas already been made (p.370) to the influence of light on tarnishing and fogging phenomena. Under immersed conditions the particular function of light appears to be the stimulation of electrochemical action. Vernon (J. Birm. Met. Soc. 1919, 7, 549) found that the illuminated areas of brass in acid solutions became anodic. On the other hand, Bannister and Rigby (J. Inst. Metals, 1936, 58, 227; J. Iron and Steel Inst. 1936, 133, 293), in a systematic investigation employing dilute solutions of potassium chloride, found that for several metals the cathodic process was stimulated, the effect in this case being dependent on the presence of

The influence of concentration of salt solutions is complex, but attention may be called to the fact that in the strongest solutions (concentrations exceeding, say, N/2) the rate of corrosion of mild steel (also zinc) falls off with increasing concentration, mainly owing to the lowered solubility of oxygen. Heyn and Bauer (1908) and Friend (ibid.) showed that in certain conditions there was a well-defined maximum or "critical" concentration, but the precise form of the corrosion rate/concentration curve may be affected by various other factors (Bengough, Lee and Wormwell, Proc. Roy. Soc. 1931, 134, A, 323; Evans and Hoar, ibid. 1932, 137, A, 343; Bengough and Wormwell, IIIrd Rep. of Corr. Ctte., Iron and Steel Inst. 1935, 148).

It will now be apparent that the rate at which a given metal can corrode in an approximately stagnant liquid depends partly on the composition of that liquid, and that this rate may vary between zero and some very high figure. Magnesium, for instance, is passive in strong caustic potash solution, but if placed in N/10sodium chloride will corrode so rapidly that it is difficult to measure the rate accurately. Iron is also passive in the alkali, but will not corrode nearly so fast as magnesium in N/10 sodium chloride. Reasons for this are the much more negative potential of magnesium, associated with its comparatively low overpotential so owing to the using up of oxygen in the system. that hydrogen can be freely evolved on the

that depolarisation due to oxidation of hydrogen hy dissolved oxygen plays no appreciable part in the corrosion process. On iron, however, hydrogen gas is not so rapidly discharged, and the rate of corrosion is largely determined by the rate of depolarisation, and this will depend on the speed of access of oxygen to the effective cathodic areas on the metal surface. This speed is determined by the convection currents and diffusion rates in the particular experimental conditions studied, and if the N/10 sodium chlorids be approximately stag nant it will he so slow that the corresponding corrosion rate will also be slow and far below that of magnesium in similar conditions. If the solution be stirred the rate of corrosion of the iron will increase with the speed of attring, but the rate of corrosion will not reach that of magnesium at any experimental speed so far tested.

An objection to the foregoing argument might he raised that owing to the rapid evolution of gas from magnesium in N/10 aodinm chloride the liquid can never he regarded as stagnant, and the comparison with iron is, therefore, imperfect. This raises the question of the effects of the motion of salt solutions on

corresion rates.

It is not easy to produce complete atagnancy in a solution since convection currents are set up by very amall temperature changes and hy evaporation, but with a minimum of motion corrosion rates may be low owing to twe factors:

Restriction of oxygen supply.
 Restriction of the supply of ions to form

solubis anodio products.

Magnesium will not be affected by (1), since oxygen plays no part in the process; it might of increased depolarisation by the rapid supply be affected by (2) if the aupply of chlorids ions of dissolved oxygen are negligible, became very low in the solution owing to the formation of insoluble hasic chlorides of magnesium. The main coreosion product is, however, magnesium hydroxide, and usually corrosion proceeds rapidly. Iron resembles magnesium in not forming stable basic chlorides, but the rate of corecesion of zinc (in a compara tively limited volume of solution) slows down owing to the diminution of chloride ions and the increase of alkalimity in the solution (weak alkalıs protect zınc, but very strong ones can attack it).

With increased motion of the liquid the rate of corrosion of iron in N/10 chloride solution also increases, at first rapidly, and then more

metal itself; this discharge of gas is so rapid | due to the formation of protective films consisting mainly of hydrated ferric oxide at all the potentially anodic areas. (In the presence of ions, such as chloride, iron will tend to travel an appreciable distance from an anode before it is precipitated as hydrated oxide), Distilled water readily shows this peculiar form of passivity, which unfortunately resembles some of the other forms of passivity in the fact that it does not persist if the metal be removed from the passivifying conditions; nor is it necessarily permanent in time even in those conditions. The protective film may gradually thicken and hreak down, and intenss local corrosion may then occur. In the absence of the passivity just described, the rate of corresion in ordinary salt solutions will increase continuously with speed of movement because of the large aupplies of oxygen available for depolarisation. Thus Speller and Kendall (Ind. Eng. Chem. 1923, 15, 134), using iron tubes which had not been previously freed from rust, found that corrosion increased continuously with increase in the velocity of

water flowing through them. The effect of motion in the case of solutions in which corrosion is mainly associated with hydrogen evolution is different from that discussed above. The rate of corresion is usually depressed as the speed increases, apparently because of the rapid removal of hydrogen bubbles which would otherwise form nuclei at which increases of gas formation could occur—in other words, gas is svolved more rapidly at an existing bubble than at a metal aurface (Whitman, Russell, Welling, and Cochrane, Ind. Eng. Chem. 1923, 15, 672) Navertheless, the rates of coreosion in such boulds are still so high that the possible effects

The effect on steel corcosion in sikali chlorides of oxygen pressures from one to twenty five atmospheres has been atudied by Les (Trans. Faraday Soc. 1932, 28, 707) and Bengough and Wormwell (IVth Report of the Corrosion Committee, Iron and Steel Inst. 1936, 213)-The effect depends largely on the experimental technique used in raising the pressure of the oxygen. If it be raised to two to five atmospheres before immersion of the steel in sodium chlorde solution (N/2 or N/10,000) or in a hard tap-water, very little corrosion occur (less than 5 mg. in 10 days on a specimen with a surface area of 15 sq. cm). If, however, the steel be first immersed in the liquid and the slowly, and this increase in, no doubt, manly pressure raised subsequently the amount of due to the increased amount of evygen made corrose in 10 days will be 120-170 mg variable for depolarisation. The effect, bear according to the solution used. If air at ever, is not the same in all selutions; for 10 atms. pressure be substituted for oxygen. instance, in some types of bard water the i.e. a partial pressure of two atmospheres of effect of slightly increased motion is to raise oxygen, the amount of corrosion in N/2 NaCl the rate of corresion, but more rapid motion will be increased to about 36 mg. in 10 days depresses it almost, or quite, to zero (Heyn and with the first method of exposure, as compared Bauer, 1910, I.c.; Friend, 1922, Ic.; Forrest, with about 90 mg. in the second method. It is Roetheli, Brown, and Cox, Ind. Eng. Chem. noteworthy that the amount of corrosion in air 1931, 23, 1010). The distribution of corrosion is less than in oxygen at equivalent partial is also gradually changed in the direction pressure. The amount of hydrogen croterior of greater localisation before complete suppressure in the direction when the second, or more usual, method of non is effected. This phenomenon seems to be exposure is used is not greatly influenced by

increase of oxygen pressure, and the corresponding amount of corrosion is negligibly small in sodium chloride solutions. The corrosion/ time curves are straight lines which show increasing slopes with increased oxygen pressure up to about 20 atmospheres; at higher pressures the corrosion rates increase but slowly, perhaps because the corrosion products become hardened and so tend to restrict the movement of oxygen and ions in solution.

It will have become evident from the above discussion that the effect of increased supplies of dissolved oxygen may he negligible, stimulative, or repressive, according to the conditions of experiment. The fact that a high supply may, in certain conditions, ennoble, render cathodic, and protect a sample of iron or steel has led to the development of the "Differential Aeration Principle of Corrosion." Early work in this field was due to J. Aston (Trans. Amer. Electrochem. Soc. 1916, 29, 449), R. J. McKay (ibid. 1922, 41, 201) and U. R. Evans (J. Inst. Metals, 1923, 30, 239), the last-named having developed the subject in many subsequent papers. The principle states that a piece of metal tends to become anodic and attacked at places on its surface where the oxygen supply is least, and cathodic and protected where the oxygen supply is greatest. An example of the principle was considered to he the behaviour of zinc or steel half-immersed in, for instance, dilute sodium chloride solution. Since the oxygen supply will he most rapid to those parts of the metal surfaces which are nearest the liquid surface, these parts should he cathodic and protected, and those lower down anodic and attacked. Many experiments showed this prediction to be confirmed. It hecame clear, however, that the composition of the liquid affected the result; thus, if ammonium chloride were substituted for sodium chloride no protection was obtained near the water-line. Accordingly, the explanation was incorporated that the protection in the solution of the sodium salt is not due to oxygen per se hut to alkali which is formed where oxygen unites with cathodic hydrogen to form water. The alkali protects the metal hy causing precipitation of ferric hydroxide on the metal at the moment iron ions enter solution. Thus the mechanism, as emphasised by Bengough and Wormwell (Proc. Roy. Soc. 1933, A, 140, 399), consists essentially in the preferential development of protective films at the most oxygenated areas of

It was next found that the protection thus afforded was not necessarily permanent and that both zine and iron could become heavily attacked at, and near, the water-line even in solutions of alkali chloride after long periods of partial immersion; in fact, the attack might bc so rapid as to overtake and surpass that which occurred at the less aerated parts of the metal. The explanation seems to be that the film which is first formed is not completely proteetive, but slowly thickens and finally breaks down owing to secular changes; when this has occurred corrosion is very rapid at the breaks, because of the high supply of oxygen which, corrosion in stagnant liquids exposed to the air ordinary temperatures. An explanation of this water-line attack which begins after long periods of partial immersion is that the supply of alkali to the water-line gradually falls owing to diminution of corrosion at more deeply immersed areas, which become covered with hardened and protective masses of corrosion product.

The differential aeration principle has been used to explain many examples of the localised corrosion of iron and steel, often with little justification. Deposits of rust, for instance, have been stated to be not only the results of corrosion hut also the causes of local attack; deposits of foreign bodies, such as sand, have been supposed to act similarly; holes and erevices in metal, to which access of oxygen is restricted, are supposed to cause pitting. None of these ideas is correct generally, though each may be true in particular conditions. Differential aeration actually appears to be only one of the factors which can influence the distribution of corrosion over the surface of a metal in particular conditions, and is often not the most important (Bengough and Wormwell, J.S.C.I. 1937, 56, 349). Others are:

1. Differences in the surface conditions of a

metal, e.g. roughnesses and phase-types.

2. The chemical composition of the electrolyte, which may become inhomogeneous when corrosion occurs.

3. The geometry of the metal-electrolyte-gas system.

Gravitational streaming.

Interfacial tension effects. Breakdown of protective films by "ageing"

and other causes.

7. The presence of foreign bodies, discontinuous mill-scale, etc.

From what has already heen said, it will have become clear that corrosion is a highly specific phenomenon both as regards metal and environment, and corrosion velocities and distribution are difficult to predict from first principles largely because of the varying physical properties of corrosion products. In a single experiment these properties may vary with time and may change, for instance, from being highly protective to completely non-protective. Hence the importance of a knowledge of long-period corrosion tests carried out in many types of defined conditions. A series of such tests has been carried out (Bengough and associates, 1926-1937) principally with zinc, iron, and carbon steels, and in stagnant conditions, but many more will be required before the subject can be considered to rest on a satisfactory quantitative basis.

Corrosion Fatigue.—Discussion so far has been confined to the behaviour of metal in an unstressed condition. If a sample of steel, for example, be stressed to a definite extent and then immersed, the corrosion rate in acids is usually increased but in salt solutions is not much altered. If, however, the stress is frequently reversed during the corrosion process a somewhat complicated process of destruction or alteration of the metal may be per se, normally acts as an accelerator of set up, the result of which is not necessarily

identical with that which would be predicted | fatigue conditions, as compared with stressless from measurements of the results of corrosion corresion, is "due to the effect of the cycle and stress sparately; it is usually much arranged atrans on the percent and rupture of the more serious. Conditions that have been particularly studied ere alternate tensons and compression, elternate bending and alternating dims are, in the writers' opinion, identical win torsion. If by any of these means the atresses are varied in a cegular manner through a range other grounds, to be commonly formed on in which zero stress is the middle point, then the steel, for instance, when corroding in salt solugreater the range the smaller the number of alterations of stress a metal can withstand before fracture. In the absence of any appreciable amount of corresion, s.e. if the experimentaire carried out in a vacuum or in a dry mert gas, a metal may be expected to withstand an infinite number of stress alterations provided the stress range is below a definite hmiting value, which is known as the "fatigue limit." Since the determination of this value is a very long and tedious process, another value, known as the "endurance limit" is often determined instead. This is merely a etress range which has not produced fracture in a stated number of stresscycles, and, as Gough (J. Inst. Metals, 1932, 49, 17) has pointed out, has no meaning unless this number is given. Gough and Sopwith (shid. 93) have shown that if teste be carried out in presence of air and moisture the curves dichromate and various emulsifying old) has showing the relation between etress range and been discussed by Burchholts and Kreketr number of cycles withstood (commonly called [Stahl u. Essen, 1933, 53, 671) and many other SN curves) he wholly below those conducted (see especially, Sopwith and Gough, J. Iron and in a vacuum or mert gas, particularly with Steel Inst. 1937, 183, 315) In the case of copper, brass, and mild steel, evidently corroson a luminum ciloys complete immunity from has played a part in the result and the joint destructive action is an example of " corresion fatigue." Much more serious deterioration may occur if a liquid corrosive agent, such as salt solution or some type of tap water, be present in addition to air. In such conditions it may be impossible to find any true " corresion fatigue limit," becouse frecture of the metal may occur at extremely low etress ranges, or even in absence of stress, i.e. by corrosson alone. The common practice is, therefore, to determine what is really on "endurance limit" at a stated number of cycles, but in much of the literature this is frequently called a "corrosion fatigue" limit.

Fracture owing to corrosion fatigue begins with the formation of surface cracks which ace narrow fissures with sharply pointed ends, and their formation and progress has been attidied in detail by Gough and associates in a whole series of papers in recent years. In aluminium, ferential corrosion along the sites of previously formed slipbands; they are numerous and amail, and are most thickly distributed "in the region of maximum resolved shear stress

tions or industrial waters. These films limit the rate of stressless corrosion in presence of a particular rate of oxygen supply, but if they were to suffer frequent rupture the corrosion rate with that same oxygen supply might be greatly increased. In this connection it is interesting to note that Speller, McCorkle, and Mumma (Proc. Amer. Soc. Test. Mat. 1928, 28, (Proc. Amer. Soc. Test. Mat. 1928, 28, 159; 1929, 29, 238) heve shown that certain passivifying agenta, such as chromates, are able to reduce liability to corrosion fatigue, doubtless by cepairing the enodic protective

film as fast as it is fractured. For steel parts operating in water, the pre vention or emelioration of corrosion fatigue by vorious surface treetments (e.g. nituding, coatings of zinc, cadmium, synthetic resins, etc) or by additions to the water (eg. sodium corresion fatigue under salt spray conditions was reported by Gerard and Sutton (J. Inst. Metals, 1935, 56, 29) from a synthetic resin varnush applied to an anodically oxidised surface of duralumin and stoved for 2 hours at 150° (see also under " Corrosion of Metals : Protective Measures") A. J. Gould (Engineering, 1936, 141, 495) has shown that the endurance of steel wire under corrosion fatigue may be conaderably reduced by rise of temperature, References - General surveys of the subject

of corrosion fatigue, with bihlographies, have been gaven by H. J. Gough [1932, supra) and by A Thum end H. Ochs, "Korrosion und Dauerfestigkert." Berlin, 1937, V.D.1. Verlag. Season - Cracking .- A phenomenon allied to corrosion fatigue is that of " season-cracking This is exhibited characteristically by fabricated articles of certain alloys, particularly high-zino brasses and nickel hrasses ("nickel silvers") following heavy working by rolling or drawing. or insufficient annealing; action may be long delayed but may cesult in the disruption of the article, sometimes whilst still in storage. As with corrosion fatigue, corrosion and atress sre the two main factors, but the latter, being static intensity; they are thus directly related to the rather than dynomic (a survival from the crystalline structure of the specimen and to the previous history of the specimen) must be epplied stress system." In general, the cracka distinguished from the alterneting stresses of do not follow the boundaries between the crystal corrosion fatigue. The corrosive agents that grains, end do not necessarily start from the promote season-crocking are usually highly bottom of large surface pits produced by corro sion. Nevertheless, atress concentration at the solutions for mild steel, and for brass traces of point of notches, etc., plays a part in the ammonia (sometimes other etmospheric impropagation of cracks when they have once been purities) or mercury; a 1% solution of mer-started. In the opinion of Gough (t.c.) the vastly curous nitrate, with 1% free nitric acid, is used increased velocity of corrosion under corrosion as a reagent for rapidly detecting season-

cracking proclivities in hrass. In further dis- ohviously a limited field of activity, since copper tinction from corrosion fatigue, season-cracks is the only one of the industrial metals that is follow an intercrystalline rather than a transcrystalline course, the internal stresses tending to draw the crystal grains apart, hence facilitating penetration of the corrosive agent; it has heen suggested that a process of "age-hardening," wherehy some constituent is rejected from solid solution at the grain houndaries, may play a contributory part. Season-cracking may he prevented by excluding the corroding medium by means of suitable coatings; much the sounder procedure, however, is to eliminate or to reduce to safe limits, hy appropriate heat-treatment, the residual stresses within the material. Much of the work on season-cracking has been carried out by H. Moore and coworkers; present knowledge of the subject is surveyed by J. D. Jevons (Metal Industry, 1937, 50, 431).

Corrosion by Sea-water.

Corrosion hy sea-water is a matter of such great industrial importance that it has received much attention in recent years. The extensive investigations of the Sea-Action Committee of the Institution of Civil Engineers ("Field Tests on Ferrous Materials," by J. N. Friend) may he especially mentioned (e.g. XVth Report, l.c., 1935). Early workers in this field believed that a solution of sodium chloride, the chief salt in sea-water, could he used for comparative tests instead of the natural water itself provided it was of equivalent (approximately half-normal) concentration. Bengough and Lee (J. Iron and Steel Inst. 1932, 123, 285) carried out experiments with the two and found marked differences. In sea-water, the corrosion of mild steel is accompanied by a much larger proportion of hydrogen evolution, which, in stagnant conditions, might correspond to one-half the total corrosion, hut in halfnormal sodium chloride solution only to ahout 5%. An increase of oxygen supply raises the corrosion rate of mild steel in sea-water hy a smaller amount than in the pure chloride solution, and this is due mainly to the fact that an increase of oxygen does not much alter the rate of hydrogen evolution. Corrosion/time curves for mild steel in stagnant sea-water fluctuate ahout a mean position and the mechanism of the corrosion is probably rather complicated. This is only to he expected, since sea-water contains, in addition to sodium chloride, chlorides and sulphates of magnesium and calcium, and traces of a large number of other salts and organic matter. Largely, no douht, on account of the last-named, it has not heen found possible to prepare from inorganic salts alone a solution which would reproduce

sufficiently susceptible to attack by ammonia. A more general vulnerability is probably offered to the action of sulphate-reducing hacteria which are very widely distributed (see "Review of the Physiology and Biochemistry of the Sulphur Bacteria," hy H. J. Bunker, Department of Scientific and Industrial Research, Chemistry Research Special Report, No. 3, 1936), hut restriction is here brought ahout hy the fact that they are active only under anærohic conditions. Particularly favourable conditions are afforded in deep-seated seamuds, as, for example, the extensive deposits in the Atlantic Ocean off Walvis Bay, South Periodic eruptions liherate large amounts of hydrogen sulphide, which not only can he detected by smell at great distances hut have heen stated to accentuate the corrosion of metal structures on land. These deposits have heen investigated by Copenhagen (Union of South Africa, Dept. Comm. and Ind., Fish and Marine Biol. Survey Div., Investigational Report No. 3, 1934), who also found the mud at the sea hottom of Capetown Docks to contain iron sulphide formed by the reaction of hacterially produced hydrogen sulphide with the excess of iron compounds present in the dock water. Corrosion associated with sulphate-reducing hacteria can he caused (1) indirectly hy the action of liherated hydrogen sulphide in solution or otherwise, or (2) more directly hy the presence of the metal structure in situ where anærohic reduction of the sulphates is taking place. In this case it is now helicved that the metal itself may take an active part in the reduction process (see "Corrosion hy Soils," infra). Examples of the first process are provided in harhours and docks, more particularly in certain waters. The presence of metals at the sea bottom must usually he due to such adventitious causes as wrecks, and interesting cases have been described where cast iron cannon, long submerged in this way, have heen completely "graphitised." Nevertheless, the possibility of a stimulation of this type of corrosion hy the presence of sulphatereducing organisms must be horne in mind under any conditions in which the supply of oxygen, locally or otherwise, is likely to become depleted.

The corrosion of marine condensers presents a number of problems that have been systematically investigated over a number of years (Reports to Corrosion Committee of Institute of Metals hy G. D. Bengough and co-workers, Corrosion is J. Inst. Metals, 1911 onwards). greater in harhour waters than on the high seas and increases with temperature and with speed

The "graphitisation" or "graphitic softening" cast iron may take place under various conditions such the corrosive effects of sea-water on mild steel.

The presence of hacteria in sca-water may also assume importance in certain cases. In the presence of nitrogenous organic matter, microorganisms of the ammonia-producing type have heen associated (Grant, Bate, and Myers, 1921) with a localised attack on hrass condenser tubes caused hy the selective solution of copper hy the action of ammonia. Such organisms have of working. Turbulence at the inlet end wall less vulnerable than, or slightly cathodic to produce rapid corrosion of brass tubes, particularly if much entangled air is prescot, giving risa to characteristic "impingement attack." Localised corrosion may also occur through the settling of various deposits in the tubes. The failure of brass tubes sometimes takes the form of so called "dezincification"; this has been shown to consist of an attack on the brass as a whole, followed by redeposition of copper in situ, and to be favoured by retention of the corrosion products as a membranous layer in contact with the metal (For ameliorative measures, see p. 389) The present position of condenser tube corrosion has been surveyed by R. May (Inst Marina Engineers, 1937, 49, 171)

CORROSION (AND PROTECTION) OF SHIPS' Hulls -The corrosion of ships' hulls comprises by far the largest scale practical problem associated with aca water attack and calls for special consideration. It will be convenient to deal here with methods of protection specific to this problem (methods of protection in general arediscussed later, p. 385) Sea-water seems to be more corrosiva than many types of fresh water towards mild atecl Thus, Montgomery and Lewis (Trans, Inst Engineers and Shiphuilders in Scotland, 1932, 75, 391) state that "vessels trading continuously in fresh water, as in the Industrial Research Council. Removal of scale Great Lakes of North America, suffar practically by acid meking has not hitherta been generally no corrosion of hull structure," Nevertheless, sea water is not necessarily more corrosive than a fresh water, for hard Teddington tap water (temporery herdness 21, permanent hardness 6 parts per 100,000) is actually more corrosive towards mild steel in stagnant conditions Movement, however, depresses the rate of corrosion of the hard water but increases that of the sea-water, so that with rapid movement the sea-water is the more corresive.

Stainless steels of the type containing about 18% chromium and 8% nickel resist the action of sea-water hetter than plain carbon ateels, hut are still too costly to use for ship con-atruction. Paints are therefore used to protect hull structures, and if care is taken to apply them properly and periodically, shell-plating, frames, and beams should last 20-25 years at least. Special parts of the ship, such as the scour area of ashes from an ejector or where a discharga of water flows more or less continuously, may require renewal in much shorter

periods.

tha plates, since a more generally distributed corresion on the plates is preferable to a localised attack on the rivets. The use of copper steel has been auggested; although the intrinsic supersority of this material is not so marked in sea water as it is in industrial atmospheres a special advantaga is claimed for its applica tion to ships' rivets, because it is alightly catholic to ordinary mild steel when the two materials are in contact

Turning to ships' plates, there is considerable evidence of changes in treatment, as distinct from changes in composition, which may have an important bearing on corrosion. It has long been customary to expose the plates to the weather, after leaving tha rolls, with the object of removing mill scale. Formerly a generous time, amounting perhaps to many months, was allotted to this process; in recent years, however, the tendency seems to have been to cut down this time and to use partially descaled ateel in the ships' hulls. It has been suggested, moreover, that the scale formed in modern practice is more difficult to remove than that formed in the older conditions; this matter is now being investigated under the auspices of the Corrosion Committee of the Iron and Steel hy acid pickling has not hitherto been generally adopted (except in the Navy), but this is a quick and efficient method, particularly if an inhibitor be used to reduce attack on the metal. The premature failure of paint costings applied to partially descaled steel is often due to the weakness at discontinuities in the scale and is aggravated by mechanical causes; when once corrosion has started, deterioration of the paint is facilitated partly by alkah produced by corrosion and partly by a process of undermining which may cause scale and paint to come away together.

Besides incomplete removal of scale, the time allowed in modern practice for painting and particularly repeinting is often quita madequate to ensure efficient protection. The dry-docking of a vessel at the end of a cruise for the purpose of repainting the hull normally occupies the period of a 12 hour tide. The operations which must be completed within this time include the removal of marine growths, the removal of old paint, the drying of the metallic surface, the application of the anticorrosiva paint and the persons.

There appears, however, to be a consensus of There appears, bowever, to be a consensus of antifolding preparation, and, finally, the drying opinion among British and Continental observers of the coating. So far as the last named process that the corrosion of both acted plates and of is concerned, tha Panti Industry has repended rivets in the structure of ships' halls has to requirements by evolving special quick drying assumed rather more serious dimensions during compositions. The older inseed oil type of recent years. In the absence of definite paints yields good results provided a sufficient evidence, changes in the quality of the metal period of drying is allowed, and combined with cannot be held responsible for the increased a red lead base it provides a priming coat corrosion of plates; but with respect to meta, that is satisfactory for general work. For corrosion of plates; out with respect to rates, that is astisfactory for general work. For many people believe that the substitution of ships' hulls, however, such pants have been "rimming" steel for abold atecl or wrought influence. (The use of rimming steel for the manufacture of rites for ships' hulls is nowing to the evaporation of a solvent, as dature or prohibited by the Committee of Lloyd's Register states the "drying" of Inneed al. In record of Shipping and certain other authorities of the proper states the "drying" of Inneed al. In record years preparations with a bituminous (asphalt) If anything, the rivets should be algebtly base have come largely into use; these have the

mediary between the steel and the antifouling paint, which frequently stimulates rather than prevents the corrosion of contiguous metal. Recent practice in the United States appears to favour the use of coal-tar and rosin, both in anticorrosive and in antifouling compositions (Gardner and Hart, Amer. Paint and Varnish Manufacturers Assoc., Circ. No. 397, 1931); the following anticorrosive is now used in the U.S. Navy Department (values in pounds per 100 gallons of paint): zinc oxide 187, venetian red 93, silica 93, rosin 146, coal-tar naphtha 383, coal-tar 48, manganese linoleate 130, beeswax 3.3. For the painting of steel hulls for fresh-water service, Speller (Mech. Eng. 1935, 57, 358) quotes good results from a mixed pigment (red lead 75%, lead chromate 12.5%, zinc chromate 12.5%) in a treated tung oil medium. J. L. Bedale (Proc. Chem. Eng. Group, S.C.I. 1935, 17, 89) gives the following as an average analysis of paint for protection of hulls in British Naval practice: 25% solvent (naptha, paraffin or white spirit), 35% soluble medium (resins and drying oils), 40% insoluble solid (zinc or iron oxide, etc.).

Assuming a satisfactory paint composition the success of the operation largely depends on prevailing atmospheric conditions; a film of moisture on the metal, even of invisible dimensions, will effectively lower the adhesion of the paint, whilst the occlusion of salt particles heneath the coating adds considerably to the difficulty. (Painting with white lead in the steel mill while the metal is still hot has heen found to he advantageous, see Fourth Report of the Corrosion Committee of the Iron and Steel Institute, 1936.) The factors that militate against the success of painting under ordinary dry dock conditions are complex enough to deserve greater time and attention than is usually accorded them if a satisfactory degree of protection is to he ensured. In practice the question resolves itself into one of economics. It is possible to secure not merely an equal but a definitely greater protection from corrosion than that obtained in earlier years; hut this is possible only by increasing the time and care expended on painting and drying with con-sequent greater cost. Which course to adopt is for engineers to decide; but it may be asked whether economy would not be better served by greater attention to protective measures, with consequent reduced wastage of metal and paint, of which latter enormous quantities are probably discharged into the sea soon after the ship leaves dry dock.

FOULING OF SHIPS' HULLS .- The "fouling" of ships' hulls by the accumulation of marine growths is not necessarily related to corrosion, for the trouble is mainly associated with the serious reduction in speed for which such growths may be responsible, and any corrosion directly attributable to this cause is usually relatively unimportant.¹ Nevertheless, it is convenient to

advantage of heing hoth quick-drying and discuss fouling together with corrosion, for the insulating, and thus provide a suitable interaccount both phenomena. The growths comprise several main groups of organisms, chiefly barnacles and molluses (both of which are commonly referred to as "shell") and certain "sea grasses" (a term which seems to be applied indiscriminately both to hydrazoa and algæ); many different species have heen recognised (see comprehensive review by J. P. Visscher, U.S. Dept. of Comm., Bur. Fisheries, Document No. 1031, 1927). As illustrating the importance of the problem, Visscher states that "after vessels have been at sea for 6 or 8 months they frequently accumulate growths from 2-3 in. in depth, and vessels with from 50 to 100 tons of fouling are seen quite frequently." Growths up to 1,000 tons in weight have been reported, but these no doubt represent cases where dry-docking has been neglected. According to Visscher, speeds in regular practice are reduced by amounts up to 50% and fuel consumption increased up to 40%. The trouble also affects flying-boats, which, in the tropics, have sometimes been prevented from rising from the water.

The treatment consists in applying over the ordinary corrosion-resisting paint a preparation that will be inimical to the attachment of marine organisms or, more particularly, their spores; such preparations are usually described as "paints," hut frequently they contain substances that would promote corrosion if not insulated from contact with the underlying metal. On the other hand, the hituminous primary coat that is commonly applied to ships' hulls, whilst protecting the metal against corrosion, offers no resistance to the lodgment of marine organisms, certain kinds of which have heen found to eat through the coating. Antifouling preparations in common use owe their value almost entirely to their poisonous content: when this is exhausted, antifouling properties cease. The problem is, therefore, to ensure that the toxic matter is released by the "paint" in sufficient concentration over a reasonably long period of time. It is one of peculiar difficulty, because widely different conditions must he met. For example, inoculation of the paint surface with the minute spores, which subsequently grow in situ to large dimensions. takes place almost entirely whilst the vessel is in port, motion being inimical to their attachment; if, however, the solubility of the paint is adjusted to afford adequate protection during this period it dissolves at an inordinate rate whilst the vessel is cruising, with the result that its toxicity is rapidly exhausted. In practice a compromise is attempted and wide variations in the effective life of the antifouling paint occur. Visscher (l.c.) states that "under optimum conditions vessels foul within 30 days of the time of dry-docking." Orton (Trans. Inst. Marine Engineers, 1931, 43, 276) questions, however, whether this figure is capable of general application (see also H. W. Keenan, J. Oil Col. Chem. Assoc. 1933, 16, 67). Light has been found to be inimical to the growth of many fouling organisms, particularly harnacles, which flourish most in "shaded" areas (Visscher, l.c.);

¹ Hedley and McLean (J. Assoc. S. African Mech. Elect. Eng. 1936, 9, 223) report, however, that stain-less steel of "18/8" type undergoes severe attack at places of attachment of barnacles.

Wolfgang Neu (Internat. Rev. ges. Hydrobiol. u. latmosphero or under water, it nevertheless wongang free inferious. Rev. ges. a problems an accordance of the Whydrogr. 1933, 28, 2283 states that green is presents problems of some magnitude indus colours of wavelengths 500-558 μμ are strongly trially. The economic importance of the avoided by eyprid larvis. The nes of suitable states, as it affects, for example, water main, coloured preparations is therefore being advo-closer cables (sheathings), and suchike, as clearly cables (sheathings), and suchike, as cated, also aluminum pigment (infru) for the reflected in the large scale organizations for its purpose of repelling growth forming organisms.

clearly depend largely on the nature of the sumlar objects has been set up in this country medium or "matrix," which should resust the under the auspices of the Institution of Civil crosive action of eca-water for at least as long Engineers and further developments must be as the toxicity of the paint is maintained; expected. various varoishee, gums, and resus are used for the purpose. Special attention is neually given to the area exposed to alternate wetting and drying (in practice a belt about 3 feet in width) moisture, and soluble matter (solid, liquid, and where mechanical conditions reach their greatest [gas], hydrogen ion concentration, electrical everity. The mixture applied to this part (the "boot topping composition") should have maximum toughness and resistance, for which purpose synthetic resins have proved especially valuable in modern practice. With regard to the toxic constituents, compounds of mercury and of copper have so far proved the most effective, their joint precence, in a fine state of subdivision, being considered to be advantageoue; pine oil is believed to add to the toxicity. M Lopez (Quim, Ind 1929, 68, 222) states that copper resinate (a green powder coluble in pain media) is preferable to most substances hitherto used, and gives results much better than those of the earlier paints. According to Keenan (lc) there is some relationship between molecular weight, effective toxicity, and structure, the cyclic aromatic organo-metallic compounds being less stable than the open-chain compounds Good results have been reported (Cardner and Hart, (c) for an anticoling preparation used (4 S C I. 183; 86, 184), attributed demanded by the U.S. Navy Department, which is as corrower attack on steel specimens embedded colleves (the per 100 gallons of panty; the (laboratory experiments) in sailed socia to a oxide 212, silica 82, magnesimm ellicate 83, cuprous oxide 112, mercuric oxide 45, balsam might reasonably be drawn from such expenroun 202, coal tar 133, pine oil 74. The presence of coal tar and of roam is noteworthy. Opinions, however, are divided as to the effectiveness of absent. Evidence is accumulating, however, coal tar in antifouling preparations and its that many examples of quite serious soil rorrouse is not generally favoured in this country, sion in service occur under amerobic conditions J. L. Bedale (I c., p. 383) states that in modern and are therefore unconnected with the supply British Naval practice the submerged part of the of oxygen. It has frequently been noted hull be given two or more coats of anticorrosive that the tendency for corrosion to take the paint, followed by one coat of antifouling comtion when exposed to air (the boot-topping composition is applied in the same way); the anti-fouling composition contains solvents similar to porous sandy soil. This has usually been those in the protective point (supra) with additions of poisonous compounds of copper, lead, arsenic or mercury. The use of chlorinated Recent investigations, however, particularly rubber as a medium, with additions of copper, by von Wolzogen Kühr and Van der Vlugt in mercury, zinc, arsenic or scienium, alone or in Holland (1934) and independently by Thaysea combination, coloured surface inimical to organisms.

Corresion by Solis.

soils is not in general so common as in the find in clay the conditions (s.e. presence of sul-

investigation, porticularly in the United States The properties of an antifording paint will and in Holland; recently a committee with

The complexity of the conditions is indicated by the large number of factors which may enter into the process, e q the content of oxygen, conductivity, partiele size, drainege; becteral processes are in many cases of importance, likewise the presence of "stray" currents from electric railways, etc , which may stimulate "electrolytic corrosion." Fortunately many of these factors are interdependent, so that conditions relatively unfavourable to corresion are by no means uncommon. Moreover, unless electrically stimulated from outside, and with the further possible exception of corresion associated with bacterial action, the rate of corresion generally tends to fall off with time because of the "self stifling" of the anodic process

Burns and Salley (Ind Eng. Chem 1930, 22, 293) found that the corresion of lead in moist, mert sands increased (within certain limits) with increasing particle size, also with mosture content of the sand and with oxygen concentration of the overlying atmosphere Bassett (JSC1, 1931, 50, 161) attributed diminished restriction of oxygen diffusion. The conclusion ments that corrosion should become negligible when oxygen is completely or almost completely form of pitting increases with the acidity of the position, the latter being applied immediately soil. It has also long been known that metal before the dock is flooded, owing to its deteriors at nectures, particularly of cast iron, are frequently more prone to undergo corrosion attributed merely to increased retention of water in the neighbourhood of the metal. has been introduced (Smith, and Bunker in this country (Chemical Research Porritt, and Dyer, B.P., 1935), aluminium Laboratory of the Department of Scientific and powder being incorporated to give a light Industrial Research), have led to the conclusion that such corrosion is in many cases attributable to the influence of sulphate reducing bacters. These are anarobic organisms of the vibro type (H. J. Bunker, Vth Rep. Corr. Ctte., Iron Although corrosion of metals embedded in and Steel Inst. 1938, 431) which very commonly

phates and exclusion of oxygen, together with | conditions little difference has been observed in the presence of minute amounts of organic matter) requisite for their activity. The organisms have the peculiar faculty of reducing sulphates, normally with liberation of hydrogen sulphide; they are typically anærobic, and while access of air does not destroy them they become active only when air is excluded. Their influence on corrosion under such conditions arises from the fact that they enable dissolved sulphates to act as "hydrogen acceptors"; an alternative mechanism is thus provided for cathodic depolarisation which is normally dependent upon the supply of oxygen. It is prohable that the process is also assisted hy a certain amount of anodic depolarisation hecause (for example) iron ions entering into solution at the anode meet with hydrogen sulphide produced at the cathode and are precipitated as insoluble iron sulphide at some distance from the anode surface. can be shown that the theoretical amount of iron converted to sulphide in this way is equal to one-quarter of the total iron corroded in the process; and in fact the presence of a certain amount of iron sulphide in the primary corrosion product is a characteristic feature of this type of anærohic corrosion. Cast iron is particularly liable under such conditions to undergo "graphitisation" (cf. deep-sea corrosion, supra) no doubt owing to the presence of graphite facilitating the expulsion of hydrogen. In practice the structures most commonly affected in this way are cast iron water mains, and here graphitisation proceeds from the outer surface (occasionally, also, from the inner, water, surface) leaving an irregular but approximately annular section of graphitised material.

Probably the most effective method of reducing the attack on iron and steel pipelines, as will be evident from what has gone before, is by attention to the method of laying. In clay soils the interposition of a layer of sand, gravel or hallast is frequently advantageous; cinders, however, should be avoided on account of the acidity that may arise from the oxidation of sulphurous impurities. Of the protective coatings applied to pipe-lines, bituminous materials, hoth coal-tar pitch and natural asphalt, are most commonly used and are frequently reinforced with felt or hessian. The extensive researches on soil corrosion carried out in the United States hy Logan and coworkers (U.S. Bur. Stands.) have heen surveyed by K. H. Logan in a number of papers (e.g. Trans. Electrochem. Soc. 1933, 64, 118; Trans. Amer. Soc. Civ. Eng. 1936, 101, 811; Bur. Stands. J. Research, 1936, 17, 181; see "Corrosion and Protection of Pipelines in U.S.A.", in "The Science of Petroleum," London, Oxford University Press, 1938, p. 747). In the corrosion of pipe-lines importance is attached to the influence of concentration cells caused by local differences, both physical and chemical, in the character of the soil in contact with the pipe; this leads to localised attack, but various other contributory factors are recognised. (For methods of soil corrosion testing, see "Symposium on Corrosion Testing Procedures," Amer.

the corrosion rates of cast iron, wrought iron, and steel (including low-alloy steels). Similarly R. M. Burns reports (Bell System Tech. J. 1 36, 15, 20) that high purity lead, lead hardened with 1% antimony, 3% tin, or 0.03% calcium, when used as cable sheathing, show approximately the same resistance to corrosion, environment heing of far greater importance than variations in the composition of the metal. Copper and highcopper alloys, however, corrode less than most of the ferrous alloys, except in soils containing hydrogen sulphide. Of metal coatings applied to the protection of iron and steel in soil, zino (hot-dipped) is most extensively used; the rate of corrosion of galvanised iron is less than that of either steel or zinc alone and when the coating is perforated zinc gives a certain amount of cathodic protection to the underlying metal. Lead, on the other hand, has a greater tendency than zinc to develop pits, and when used as a coating tends to accentuate, if perforated, corrosion of the contiguous iron. (See also "Corrosion of Metals: Protective Measures," e.g. bituminous coatings, p. 395).

G. D. B. and W. H. J. V. OF METALS: PRO-CORROSION TECTIVE MEASURES. Methods of com. bating corrosion may be considered as falling into three groups according as the result is achieved by (1) the control of external factors, (2) the use of corrosion-resistant alloys, (3) the use of protective coatings.

(1) Control of External Factors.

Methods of protection depending on the control of external factors differ considerably in detail. Frequently they consist in the removal of some harmful constituent from or the addition of inhibitive material to the corrosive medium. It follows from what has gone before that corrosive atmospheres can he rendered inocuous either by filtering out solid pollution or by suitably absorbing gaseous pollution; but such measures can evidently be applied only in very special cases. A possible and ohviously efficacious method of approach is to remove the pollution at source and much progress is being made in this direction. In the light of the principle of critical humidity, corrosion can frequently be avoided by adjusting the relative humidity of the atmosphere so that it is maintained consistently helow the critical value. A knowledge of general principles is particularly important, since it is clear that the corrosion mechanism must he understood hefore the appropriate kind of "hrake" can he devised.

In the case of immersed corrosion, it is again evident that methods involving treatment of the corroding medium cannot in general be applied on the large scale; hut on limited scales and more particularly in enclosed systems such methods are, in fact, of great importance. The procedure to he adopted will depend upon whether corrosion is mainly of the hydrogen evolution or oxygen absorption type. As discussed in the general article the hydrogen evolution type of corrosion is encouraged Soc. Testing Materials, 1937, 95). Under soil by low $p_{\rm H}$. Certain acid types of natural

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by suitable adjustment of p_0 , usually by neutral solutions under many service conditions addition of hims, which in effect displaces corrosion from the hydrogen evolution to the alone by the removal of oxygen from the system. oxygen absorption range, the rate of attack being at the same time very greatly reduced (Whitman, Russell, and Altien, Ind. Eng. Chem. 1924, 16, 665).

In cases where corresion is inevitably of the hydrogen evolution type, use may be made of overpotential. A familiar example is the amsignmating of zinc electrodes for the prevention of local attack by the acid electrolyte; here the normal process, which would result in the liberation of hydrogen from the zine surface, is suppressed because of the high overpotential of hydrogen in respect to mercury. An interesting practical application occurs in the acid pickhing of iron and steel articles for the removal of surface oxide (scale and rust). Attack of the acid upon the metal, as distinct from the oxide, involves the liberation of hydrogen; hence this part of the process can be suppressed hy such additions as will increase the hydrogen overpotential For this purpose organic in-hibitors are used, of which a varied choice is available, beterocyclic nitrogenous substances (e.g. pyridine) being particularly effications. Mann, Lauer, and Hultin (Ind Eng. Chem. 1936, 28, 159) examined the inhibitive effect of ali phatic amines on the acid attack of iron and concluded that the amount of surface blanketed by any given molecule (henco the efficiency of the inhibitor) is dapendent on the length of the hydrocarbon chains, the nitrogen stom being hald responsible for the actual attachment to the metal. Many proprietary inhibitors used in picking iron and steel are derived from by products of industrial operations, especially coal-tar, W. Machu (Korrosson u. Metallich, 1937, 13, 20) in a comprehensive survey, suggests that the inhibiting action at the cathode is due to the formation of a film baving high electrical resistance.

An application of the overpotential principle occurs in connection with magnesium-rich alloys, the corresion of which invariably follows the hydrogen evolution type. One of the uses of the anodic process may result in its localisaproposed for these alloys is the manufacture of fuel tanks for aircraft, but difficulties arise been emphasized by Evans (Trans, Electrochem from the introduction of "leaded firels" in 500, 1933, 69, 213).) Solution silicate has been anti knock purposes. The lead is used in the used for the treatment of water intended to pass form of lead tetra-ethyl, and ethylene dibromide through iron and steel systems, but its success is added for the purpose of removing the lead in the exhaust gases. In the presence of water a complicated series of reactions causes deposition of part of the lead on the surface of the alloy which then becomes attacked with liberature of hydrogen. The organic inhibitor process consists in mixing with the fuel a small quantity of a petrol soluble substance that will suppress of a petroleoluble nustance was two representations of a petroleoluble nustance was the evolution of hydrogen from the alloy produce attend of about 1%, has been found to be extent of about 1%, has been found to be extented about 1%, has been found to be extended to the number of the full are unproof "when it contains desolved ailes. impaired (DSI.R. Rep. Chem. Res. Board, 1934-7, London, H.M.S.O., 1938).

water may have their aggressiveness removed oxygen, as commonly obtains with waters and Fur this purpose both chemical and mechanical methods are employed. The former consist usually in the passage of water (which for econumical working must be heated) over a large aurface of iron, special precautions being taken to prevent the choking of the system by the rust produced. Suitable plant is described hy Speller (" Corrosion, Causes and Prevention," 2nd ed., 1935). Other chemicals, such as sodium sulphite, are assertimes used for the removal of oxygen, but considerable care is necessary to avoid excess and the content of dissolved salt is in any case increased. There are many types of mechanical deserators, from open heaters, in which a current of steam sweeps over the surface of the heated water, to heaten of the Elliutt type, in which water is introduced into a chamber at reduced pressure, the sudden boiling on superheating serving to expel dissolved gases. For the deacration of water in the cold a fine spray is injected into a chamber under high vacuum. Mechanical descration has the advantage of removing carbon dioxide and other dissolved gases besides oxygen, but complete elimination of the latter is probably best achieved by means of a supplementary chemical treatment.

Further methods of water treatment include the addition of substances that will (1) suppress the anodic as distinct from the cathodic process which is the function of the "inhibitors" previously considered, (2) lead to the formation of a film or scale over the whole surface. The first group comprises alkalı additions (sodium hydroxide, sodium carbonato, or calcium bydroxide); these become less effectiva with ancreaso an concentration of dissolved salts, particularly chlorides. It also includes chrom stes, which are frequently added to brine solu tions for corrosion prevention. In either case such additions must be made with caution, for if, as as most often the case, the rate of corresion is under cathodic control, incomplete suppression tion, with consequent pitting, (This point has been emphasised by Evans (Trans, Electrochem appears to he dependent on the presence of calcium and magnesium salts, which assist in the deposition of a protective coating. In the case uf a salue water it must be regarded in the same category as the alkalis (supra) and appropriate care exercised in its addition. Nevertheless, on lead, in various types of water, silicate additions of the order of about 10 parts

Organie colloids such as gelatine, agar, etc. also exert considerable protective influence in In the case of corresion processes the rate of neutral solutions (Friend and Vallance, JCS. which is controlled by the amount of dissolved 1921, 121, 460), the mechanism consisting phase on the metal surface (such emulsoids are known to retard solvent action of various kinds when adsorbed at the solid/liquid interface). The comparatively long life of metal cooking utensils (particularly of alumininm) has been attributed to the colloids present in food. Patterson and Culbert (J.S.C.I. 1935, 54, 327T) have described another type of inhibitor, represented by the restraining effect of polyhydric alcohols on the corrosion of iron in potassium sulphate solutions. Adsorption of the organic substance on the corrosion product was believed to be responsible for subsequent retardation, which was found to depend upon the number of hydroxyl groups per molecule of inhibitor.

Hard waters are in general less corrosive than soft waters because of their ability to deposit surface films that are protective in character. Provided carbon dioxide is not present in excess, incipient corrosion will often favour the precipitation of films by reason of cathodically produced alkali. In one successful method of water treatment for pipc-lines, just sufficient alkali is added to give a protective layer over the interior of the pipe. Very soft waters are sometimes treated with limestone in order to introduce film-forming constituents. The principles underlying the artificial treatment of natural waters for the controlled precipitation of suitable protective coatings have been of suitable protective coatings have been developed mainly by Tillmans in Germany (e.g. Gesund. Ing. 1912, 35, 669; Gas u. Wasserfach. 1931, 74, 1097 seq.) and by Baylis in America (e.g. J. Amer. Waterworks Assoc. 1935, 27, 220). In each case the primary object was the protection of the inner surfaces of supply pipes and the conditions sought were such as would produce a layer of calcium carhonate, or a Tillmixture of calcium carbonate with rust. mans' work was based mainly on the conception of the aggressive action toward calcium carbonate of waters containing more than a certain amount of free carbon dioxide; i.e. part of this carbon dioxide is required to keep bicarbonate in solution, hence it is only excess above this amount that enables water to attack calcium carbonate, so preventing the formation of a protective layer on the iron. The excess was called by Tillmans "aggressive carbon dioxide," a term which, it should be observed, refers only to aggressiveness towards deposits of calcium carbonate and not towards metal, although that may follow indirectly. (The direct action on iron, at ordinary temperatures, of such concentrations of carbon dioxide that are likely to be present in natural waters is very much smaller than was previously supposed and is probably negligible in most instances, although it may bccome significant at higher temperatures as in closed heating systems.) Aggressive carbon dioxide may be removed by contact with marble or limestone. Baylis (l.c.) has described examples of the successful application of these principles and has given curves connecting the determined $p_{\rm H}$ of the water with equilibrium values for alkalinity in terms of calcium carbonate. If

presumably in adsorption of the disperse; to precipitate out from the water and form a coating, whilst if helow the curve the water is "aggressive" and tends to dissolve calcium carbonate. Baylis emphasises that the curve is modified by other constituents, magnesium carbonate, etc., and it is necessary to determine the calcium carbonate equilibrium point for each particular water hy direct experiment. Lime is usually used to treat the water but soda can he used if it is desired to avoid an increase in hardness.

Problems connected with "external factors" under soil conditions are discussed in the

general article (p. 384).

In power plants and in many industrial operations it is often necessary to use water from which scale-forming constituents have been removed in a softening process. Generally the need is dictated by objections arising from accumulations of deposits; in boilers, however, the presence of magnesium salts is deleterious from the corrosion point of view and their removal therefore becomes important. For considered, already however. reasons softened water should be rendered slightly alkaline; small amounts of certain other materials are also commonly added for the purpose of producing protective films. is needed in using waters that have been softened by base-exchange processes hecause of their higher content of dissolved sodium salts; contact of dissimilar metals, e.g. steel with copper or hrass, should be especially avoided. A method of ameliorating the corrosivity of such waters is to hy-pass a certain amount of the raw water so that the treated water may have

the necessary minimum degrees of hardness.

Corrosion in refrigerating systems, which affords good examples of protection hy control of external factors, has formed the subject of a general discussion (Proc. Brit. Assoc. Refrigeration, 1936-7, 33, 35). Corrosion may affect either the ammonia (refrigerant) circuit or the calcium chloride (brine) circuit, but appreciably so only in the presence of water in the refrigerant or of oxygen in the hrine. A third possible contingency is the leakage of refrigerant to the brine; this must be especially avoided because of the formation of ammonium chloride, hy which the hydrogen evolution (as distinct from oxygen.

absorption) type of attack is actively promoted. Electrochemical Methods of Control.—Corrosion in aqueous solutions and natural waters being an electrochemical process (attack on the metal proceeding from anodic points or areas) "electro-chemical" or "electrolytic" methods aim at countering the natural process by rendering the whole metal surface cathodic, for which purpose an external electromotive force is employed. The principle is attractive and its application can be readily demonstrated by small scale experiments. Difficulties arise, however, on the large scale owing to the impossibility of distributing the current uniformly over the whole of a large area. For example, in condenser systems (in which electrolytic methods, at one time extensively tried, have heen largely discontinued) anodes were inserted in the water for any given water the determined $p_{\rm H}$ falls boxes at the ends of the tunes, which themselves above this curve, calcium carbonate will tend constituted the cathodes; in practice, however,

no more than the first few inches of the tubes so-called "stainless irons") to about 0 3% as could hope to receive protection (Bengough and in stainless cutlery, and methods of heat-May, J. Inst. Metals, 1924, 32, 245). Neverthe-treatment have been improved. There is conless, for the protection of pipe-lines in soil, where siderable evidence that the stainless properties the function of an electrolytic method is to enpplement the protection afforded by the usual protective coating, successful application has is self-healing, provided that environmental been claimed, Examples are quoted by K. H. Logan (I.c. "Science of Petrolenm," 1938).

A method similar in principle to that of an externally-applied E M.F. is the use of protector blocks, as, for example, the zinc alabs attached to hronze propellor hlades and to the adjacent steel hull for the purpose of deflecting electrochemical attack from the steel atructure to the readily-replaceable zinc. The method is limited hy the (normally) very short distance to which the cathodic protection can extend, to meet which objection a series of such slabe are usually fitted. Much greater effective distances for the protection of duralumin by zinc were found by Akimov (Korrosion u. Metallich 1930. 6. 84) and zinc pistes are in fact used for the protection of the duralumin portious of wing tip floats of aircraft (W. H. Hatfield, J. Roy. Aeron Soc. 1935, 39, 564). In closed systems zine may operate by using up available oxygen, for avample, Bannister and Kerr (Trans. Liverpool Eng Soc. 1933, 54, 15) reported equally good protection from zine blocks attached to steel boiler plates when electrical contact was prevented, showing that in this case protection was chemical rather than electrochemical.

(2) Corrosion Resistant Alloys.

It was formerly considered that increase in the cerrosion resistance of a metal could be effected only by more completo purification or by additions such as would render it more "noble" or intrinsically more resistant. Much more commonly in modern practice the result is achieved by additions that are themselves intrinsically more resctive, the rapidly formed initial film imposing increased resistance to subsequent attacl. Much depends on environment, and an alloy that will develop a highly resistant film under one set of conditions may fail to do so when the conditions are changed, a fact that must be horse in mind in the selection

of materials for various purposes.

Iron-rich Alloys (Stainless Steels, etc.) — Industrial non-rusting steels fall into two main groups: (1) Chromium ateels, represented typically by ordinary "atainless steel" containing 13-14% chromium; these are capable of being hardened by quenching from a suitable temperature. (2) Austenitic nickel-chrommm steels containing typically 18% chrominm, 8% nickel; these alloys cannot be hardened by openching. A third type, more recently observed in industrial atmospheres high in introduced, contains 16-20% chrominm and aniphur dioxide. Rusting is not prevented, but about 2% nickel. As this is non-anstemment at it is less destructive in character. Moreover, resembles the first group in its shilty to harden, such atcels provide an improved bend both for resembles to mit group in its samp violation; since after properties it is plant contings and improve bend tool whit in corrosion-resisting properties it is plant contings and for galvaniang. Chromizometeroliate between groups (i) and (2). The opper mild steels, containing approximately typical alloy of the first group is simular in [18] chromizm, 0.5% copper, are stated to show chromizm content to Brearley's original masterial improved corrosion-resisting properties over (1013), but carbon now ranges from 0.1% (the ordinary copper steels (see p. 34).

conferred by chromium are due to the presence of a teus crous oxide film, rich in chromium, that conditions are sufficiently oxidising, and prowided also that the surface is smooth-preferably " hright polished." Ordinary stainless steel is highly resistant to tap-water, fruit juices, and all oxidising acids (contact with strong nitric and actually strengthens the passive film and improves subsequent resistance), but is readily attacked by non-oxidising acids (e.g. hydrochlone and sulphuric), hy which the passive film me destroyed. Particles of chlorides as in manne atmospheres and of soot in industrial atmospheres are immical, and the auriace should therefore be wiped frequently under such conditions In sea-water ita resistance is onl moderate and it is lishle to snodic strack when in contact with non-ferrous materials.

The presence of nickel in the "18-8" austenitie alloy, coupled with the fact that the added elements are in solid solution, gives a greater range of stability to the invisible passive film, and in particular a greatly improved resistance to sca-water with freedom from electrolytic corrosion in contact with ordinary steels and non ferrous materials. The alloy has also an improved resistance to sulphune acid and, although to a much less extent hydrochlome acid; further resistance is obtained by the addition of molybdenum (typically 25%) and certain other elements. Austention stainless steels are especially resistant to oxide. tion at high temperatures, but if they are memtained for any length of time at temperatures between 500° and 900° (as in welding) they are hable subsequently to an intercryetalline type of corrosion (" weld decay ") due to the partial rejection of chromium carbide to the grain boundaries. The condition can be removed by appropriate heat-treatment or, alternatively, may be prevented by the addition of small amounts of certain elements (e.g. titanium) which have the effect of retsining carbides in solution; nichium (about 10 times carbon content) being less easily oxidised than titanium) is recommended for steels that bave to withstand the most drastic welding treatment. lleat resisting etecls bave heen surveyed, with extensive hibliography, by W. H. Hatfield, J. Inst Fuel, 1938, 11. 245.

For large-scale structural work, for which atamless steels would be too costly, copper ateels, containing 0 25 to 0 5% Cu, are available The increased resistance to corrosion (which has been identified with the formation of copper-rich surface film) is more particularly High silicon cast irons containing from 13 to 15% silicon ("duriron," "tantiron," etc.) are much used in chemical engineering for acid, pnmps, etc. Their resistance to sulphuric and nitric acids is very high. A similar alloy ("durichlor") containing the same amount of silicon with 3.5% molyhdenum is stated to he especially resistant to hydrochloric acid, its relative immunity heing very prohably due to the formation of an insoluble coating of molyhdenum chloride. (For handling hydrochloric acid in chemical engineering practice it is, however, generally advisable not to depend upon resistant alloys hut to use where possible either glass-lined or ruhher-lined equipment.) Austenitic cast irons, containing typically 14% nickel, 6% copper, 2% chromium, have, hesides greatly improved mechanical properties, a high resistance to many kinds of chemical attack, comparable with and sometimes superior to that of phosphor hronze (J. G. Pearce, Proc. Chem. Eng. Gp., S.C.I. 1934, 16, 91; A. B. Everest, ibid. 1937, 19, 69). They have also high resistance to oxidation hy heat, with freedom from "growth," to which ordinary cast irons are liable after repeated reheatings.

Nickel-rich Alloys.—The high resistance exhibited hy nickel towards many industrial liquids, particularly alkalis, is shared markedly by its principal alloys. Of these, monel metal (a "natural" alloy of nickel and copper smelted direct from the mixed ore) is the most typical and most widely used. It contains approximately 68% nickel, 28% copper, 2% iron, remainder manganese, silicon, etc. Similar alloys are prepared synthetically. They are characterised by generally good resistance to non-oxidising acids, e.g. monel metal is widely used for pickling equipment using hot dilute sulphuric acid. It is also very suitable for superheated steam fittings, pumps, etc. Monel metal has only a fair resistance to hydrochloric acid (although hetter than either nickel or copper alone); greater resistance is obtained by the substitution of molyhdenum for copper, and this is the hasis of certain proprietary alloys recommended for use with hydrochloric acid. All the "non-oxidising acids" have their corrosivity greatly increased hy aeration. This is shown by the following figures, due to Thompson and McKay (Ind. Eng. Chem. 1923, 15, 1114) (for specimens moving at 15 ft. per min. immersed in dilute sulphuric acid at 82°C.):

Corrosion Rate.

Mg. per sq. dm. per day.

Air-free (washed with methane).

Copper . 43 1,070

Monel metal 64 700

The advantage of an "air-healing" film as in 18/8 chromium-nickel steel is shown by the further data (Searle and Worthington) quoted by Moore and Liddiard (Chem. and Ind. 1935, 54, 787).

Loss of wgt., mg. sq. dm. per day, in 3 % sulphurie acid: Oxygen- With oxygen free. bubbled through.

Monel metal . 17 630 18/8 CrNi steel 442 4 Alloys of nickel with chromium (see also in General article under High-temperature Oxidation) are used in dairy and other food industries and in chemical engineering, where high corrosion-resistance is required. "Inconel" contains approximately 80% nickel, 14% chromium, 6% iron. Nickel-rich (chromium-free) alloys are not good for oxidising acids (e.g. nitric acid); they have, however, extremely high resistance (not appreciably reduced by aeration) towards alkaline solutions, caustic soda and caustic potash. Aqueous ammonia is resisted if dilute, but stronger solutions are aggressive, particularly towards nickel-copper alloys. (For such conditions ordinary iron or steel, which is completely passivated hy ammonia, may he used). Nickel alloys have a very high resistance towards sea-water (see "copper-rich alloys"). They are sensitive to attack by sulphurous acid solutions and hy humid atmospheres containing traces of sulphur dioxide.

Copper-Rich Alloys.—Copper-rich alloys fall into two main groups, (1) copper-zinc alloys or hrasses, (2) alloys of copper with other metals—principally with tin (the true hronzes), with aluminium (aluminium hronze), and with nickel (cupro-nickel). Of these the corrosion-resistance of the first group is generally less than that of copper, whilst members of the second

that of copper, whilst members of the second group are in this respect at least equal to and frequently superior to copper. Whilst the hrasses, therefore, are generally selected primarily for economic or mechanical considerations, additions heing made where necessary for increasing resistance to corrosion, the resistance of the other alloys is frequently of major importance and, together with their

superior mechanical properties, dictate their application in chemical engineering.

Brass is used extensively as tubes, etc., e.g. in evaporators and in condensers. Under certain conditions, particularly where insoluble corrosion product can accumulate in contact with the metal, as, for example, in marine con-denser tuhes, so-called "dezincification" may occur. The net result is a removal of zinc and enrichment of copper, hut the process has been shown to consist in attack on the alloy as a whole followed by redeposition of copper, with consequent weakening of the material. Dezincification is more prone to occur in 60/40 hrass which contains crystals of α and β solid solutions (the β zinc-rich constituent undergoing preferential attack) than in 70/30 hrass which consists entirely of the a phase. The addition of 1% tin as in the Admiralty (70-29-1) alloy confers rather greater resistance, whilst com-plete immunity from dezincification is obtained under many conditions by the addition of 0.01% arsenic. Another type of rapid attack caused hy the impingement of an aerated stream of water ("impingement attack") has heen overcome by the addition of 2% of aluminium, and this material (containing 76% copper, 22% zinc) is now widely used for condenser tubes; with the further addition of 0.02% arsenic for the suppression of dezincification, it has largely superseded Admiralty hrass for condenser tubes. (For further information on marine condenser tuhes, see references on pp. 381, 382.)

The copper-tin alloys find extensive applica- | are "Mg7" and "Hydronshum" (approxition for pumps, valves, etc. and, with the mately 7% magnesium). The addition of small addition of a little zinc (eg. Admiralty gun- amounts of antimony as in the German "KS. metal, 88 copper, 10 tm, 2 mel, for various Seewasser alloy is claimed actually to increase metal, and the fittings. Phosphor brones as a life of formation of anilmony mychloride) the copper-tim alloy to which a small amount of resistance of the natural film to seewater. In phosphorus, up to 05%, has been added as a deoxidiser. They have good general corrosion resisting properties. The aluminum bronzes (copper with 5 to 10% Al), owing to the formation of a surface film of aluminum oxide, are useful in many conditions, as, for example, in contect with hot and cold sulphuric acid in moderate strengths; the alumina film also enables them to resust oxidation at high temperatures. The copper-rich nielel alloys atrongly resemble the corresponding nickel-rich alloys in resistance to sea-water corrosion particularly impingement attack (supra), and cupro-nickel tubes, approximately 70% copper, 30% nickel, are also widely used in marina condensers.

All the copper-rich alloys are vulnerable to trio acid. Their response to non-oxidising nitrio acid. acida depends in general upon whether or not the solution is aerated In the absence of air, high resistance is shown, but severe attack may occur if air is admitted (of behaviour of copper, Thompson and McKay, supra). Resistance to caustio alkali colutions is good, but not so good as that of the nickel-rich alloys. The copper rich alloys resemble copper in susceptibility to attack by aqueous animonia and their use should therefore he avoided in the presence of this reagent. The use of copper alloys in chemical engineering bas been surveyed by S. Baker (Proc. Chem. Eng. Gp. S.C.I., 1937, 19, 41).

Aluminium and Magnesium Alloys -Whilst the use of aluminium and magnesium alloys is most commonly dictated by considerations of lightness and "atrength-weight ratio," their hightness and resistance to corrosion may assume great importance, as, for example, in aircraft exposed to see apray. Aluminium differs from the metals previously discussed in that its highest corrosion resistance is characteristically exhibited in the unalloyed state; this is becausa of the continuity of tha thin covering film of oxide to which the metal owes its normally good corrosion resistance, notwithstanding its intrinsically reactive character. Recent develop-ments in the industrial electrolytic refinement of ments in the industrial electrolytic refinement of contradistanction to aluminium, with a relatively aluminium have led to a product in highest low "intrinsio" reactivity, like aluminium, purity (exceeding 99 99%) from which extra Investre, the freshly-cleaned metal become ordinarily high resistance to chemical attock by rapidly covered with a film of oxide which has acids and hy aca-water has been reported (R. marked protectiva properties (Vernon, Trans Gadeau, Chem. et Ind. 1935, 34, 1021). Certain Faraday Soc. 1927, 23, 156). In contact with of the aluminium alloys having maximum sulphuric acid this film is converted to relatively

alloys of the duralumin typa the presence of copper confers valuable age-hardening pro-perties, but such alloys, if the necessary beattreatment has been imperfectly conducted, sre hable to undergo intercrystalline corroson, particularly in marine atmospheres. Aluminium and its alloys are used over a wide range of conditions, atructural, architectural, industrial, and domestic. For cooking utensils they have the advantage of complete non toxity. to the solubility of aluminium in alkalies, skaline conditions should in general be avoided; for the cleaning of aluminum utensils ordinary sods should be replaced by preparations containing sodium silicate which is much less harmful,

The greater lightness and favourable strengthweight ratios of magnesium base alloys (eg. "elektron" types ranging from 90 to 96% magnesium, with additions mainly of aluminium, manganese, and zine) have encouraged their use within recent years. The readiness, however, with which magnesium oxidises superficially is not offset, as with aluminium, by the resistance of the oxide film towards marine atmospheres; moreover, attempts to produce a protective type of oxide film by anodic processes, as for aluminsum, bave so far proved unsuccessful. Other protective coatings are available (see "Protection by Coatings," infra) that will resist severe salt-apray conditions, but at present there is considerable reluctance to displace the usual aluminium alloys for aircraft, although in other directions the use of magnesium alloys appears to be extending. Magnesium and magnesium-base alloya bave extraordinary resistance to hydrofluoric soid (through formation of an insoluble film of magnesium fluoride) and also (in contrast with aluminium) to alkalis and alkalina solutions. Other data are given by L. Whitby (1nd. Chem. 1931, 10, 203). Alloys of the light metals have heen surveyed by C. H Deach (Froc. Chem. Eng. Group, S C.I. 1937, 19,

Lead Alloys.-The corrosion resistance of lead, which under many conditions is the main enterion determining its use, is associated, in of the aluminium alloys having maximum sulphures acid this film is converted to relatively another than the presence value in a strength, durations, it is given persona value mounted in the presence of the matural protective film; such alloys are best treated by the another ordinary to the natural protective film; such alloys are best treated by the another ordinary in the protection of process or by a "metal-cladding process" (see the sudder "Protection by Coatings," in first, a line of the protection will obtain a new much less than 100 miles of the protection o for atmospheric and sea air conditions; examples the resistance of lead, and also of lead antimopy

lead" alloys contain usually from 10 to 15% antimony; they are much used in chemical plant, particularly for acid pumps. Alloyed with 10 to 25% tin, lead is used as a covering for iron ("terne plate") for roofing purposes (see p. 374). For cahle sheathing, water-pipes, etc., lead is usually strengthened by certain additions, e.g. small amounts of antimony or tin. or traces of calcium; high-strength ternary alloys, containing cadmium (0.25%) with either tin (1.5%) or antimony (0.5%), and characterised hy freedom from intercrystalline failure to which pure lead is liable under conditions of vihration, have been developed in this country (S. J. Nightingale, Dev. Rept. D.3, Brit. Non-ferrous Met. Res. Assoc. 1931). None of these additions, however, appears to affect appreciably the corrosion-resistance under soil conditions (see "Corrosion hy Soils," p. 384) A similar conclusion probably holds good for most water conditions also; see, for example, Nightingale, ibid.; F. Mayer, "Wiener Hochquellwasser und Bleirohre," Berlin, 1934. Nevertheless, the susceptibility of lead to corrosion in soils containing decaying organic matter, and the "plumbo-solvency" of lead in certain natural waters must be horne in mind. The behaviour of lead and lead-base alloys towards natural waters has been surveyed, with extensive bihliography, by H. Ingleson (D.S.I.R. Water Poll. Res. Tech. Paper No. 4, London, H.M.S.O., 1934). Reference has already been made (p. 374) to the need for avoiding contact with oak; it is also desirable to insulate lead (e.g. by bitumen impregnated felt) from contact with cement, which, in presence of moisture, may cause serious corrosion (D.S.I.R. Building Res. Bull. No. 6 (3rd ed.), London, H.M.S.O., 1935).

Tin Alloys.—For economic and other reasons tin is most widely used in the form of coatings on iron and copper, tinplate as used in the canning industry heing the outstanding representative; familiar exceptions are tinfoil for the wrapping of foodstuffs, block tin for condensers (distilled water) and for the handling of heer, sheet tin for collapsible tubes (dental pastes, etc.). The principle tin-rich alloys are Britannia metal and powter; formerly Britannia metal contained from 5 to 8% antimony, and pewter from 8 to 15% lead. Modern pewter, however, contains typically 5% antimony, 1 to 3% copper, and no lead. From the corrosion-resisting stand-point T. P. Hoar (J. Inst. Metals, 1934, 55, 135) has shown that the presence of 5% antimony is desirable, but suggests that the further addition of 1 to 3% copper is deleterious.

Silver Alloys.—The application of silver and silver-rich alloys in chemical engineering is naturally limited hy economic considerations, hut a number of practical uses, particularly for resistance to attack hy alkalis, both fused and in solution, has been described by D. McDonald (Proc. Chem. Eng. Group, S.C.I. 1931, 13, 50). "Sterling silver" (92.5% silver, 7.5% copper)

alloys, to sulphuric acid (Singleton and Jones, action of traces of sulphur compounds in the J. Inst. Metals, 1933, 51, 71). Type B "hard atmosphere. Reflectivity measurements (Veratmosphere. Reflectivity measurements (Vernon, Trans. Faraday Soc. 1924, 19, 882) showed a reduced amount of tarnishing on sterling silver containing 1.75% cadmium, provided that the surface was hrightly polished. H. A. Sloman (J. Inst. Metals, 1934, 54, 161) found that the addition of as little as 0.5% heryllium to silvercopper alloys approximating in composition t) sterling silver conferred a marked degree of inmunity to tarnishing under quite drastio conditions, owing to the formation of a protective self-healing film of heryllia.

Zinc Alloys.—Although alloys in which zinc is the main constituent have little application as corrosion-resisting alloys there are cases in which corrosion becomes an important factor in determining their use. An example is that of the zinc-hase die-casting alloys (approximately 4% aluminium) which, unless special attention to a form of intercrystalline corrosion, particularly under warm, humid conditions. The action is accompanied by a dimensional change or "growth" (for discussion, see Russell, Goodrich, and Cross, J. Inst. Metals, 1928, 40, 239); this growth is probably caused hy oxidation of zinc at the grain houndaries, hence the high oxide/metal volume ratio of zinc (p. 369) would be expected to contribute to mechanical failure. Minute traces of impurity (particularly lead) are primarily responsible and no trouble whatever is experienced when zinc of the highest purity (exceeding 99.99%) is employed. For zine used as a protective covering to iron under urban atmospheric conditions the presence of lead is advantageous (W. S. Patterson, J.S.C.I. 1927, 46, 392) no doubt owing to the accumulation of relatively insoluble lead sulphate in the product.

References (Books dealing with Corrosion-resistant Alloys): J. H. G. Monypenny, "Stainless Iron and Steel," 2nd ed., 1931, Chapman and Hall, London; McKay and Worthington, "Corrosion Resistance of Mctals and Alloys," 1936, Reinhold Publishing Corporation, New York, See also works recent poration, New York. See also works recommended under Corrosion, General (p. 368). Moore and Liddiard, "Corrosion-resisting Nonferrous Alloys," J.S.C.I. 1935, 54, 786.

(3) Protection by Coatings.

Protective coatings may he classified according as they have or have not a chemical bond with the underlying metal. The former include hoth "natural" and artificially stimulated coatings of oxide and certain corrosion products. The latter comprise chiefly (a) paints, varnishes, and such-like, (b) metal coatings.

Natural Protective Coatings.—A number of commonly used metals have appreciable resistance to corrosion hy virtue of a film of oxide with which they are normally covered. Aluminium, for example, has a heat of oxidation among the highest of the common metals and yet is comparatively stable under many conditions. The formation of invisible oxide films widely used for silver plate and ornamental work, on iron and copper has already heen discussed; is prone to discolour hy tarnishing through the "stainlessness" of stainless steel is due to

the presence of e tenecious oxide film that is self-| chromium oxide) has good resistance to wear as heeling under slightly oxidising conditions. Such invisible protective films may not inoppropriately be called "natural varnishes" A feeture common to them all is that the protection breaks down, sometimes spectacularly, if the external conditions after in such a way that the oxide film cennot be meintained.

A good example of a "natural paint" is provided by the green patina on copper. This is e genuine corrosion product and is characteristic of attack by our charged either with ealt sprey at the ceabeard or with traces of combustion products inland, the essential constituent being basic copper chloride or basic copper sulphate, the latter, however, being much the more widely distributed (see under General article). The protective character of the patina is probably due lergely to the continuity and insolubility of these basic com pounds, although there is some evidence that a thin film of cuprous oxide, which is invariably found beneath the green deposit in immediate contact with the metal, plays an important part Unfortunately, it is the exception for a naturel corresion product to behave in this way, frequently its presence encourages further attack on the underlying metal.

Artificial Coatings Reinforcing or Simu-Protective coatings of oxide may be produced by thermal, chemical, or electrochemical treatment, An old-established example of the thermal method is that of the Bower-Barff process for gron and ateel, with its various modifications (Gesner, Bontempi, etc.) the essential part of which consists of heating the metal with superheated steam. Copper and copper rich alloya heated steam. Copper and copper and empa-may he protected against tarnishing by thin films of oxide produced by low-temperature heating in air (Vernon, J.C.S. 1926, 2273); these may either retain essentially the original appearance of the metal or, according to temperature and time of heating, give pleasing interference colours that are highly permanent as distinct from films of sulphide of similar initial appearance. Immersion in on oxidising liquid high current density) a green patina is produced. so sometimes employed, es, for example, the which, however, at first consists of a base "chromato" dip for magnessum and the esulphate, CuSO_Cu(OH), analogous with MB.V. [Modified Bouer-Vogel) process of saladicute. On subsequent exposure, the colour aluminium; but such coatings here the disadvantage that they are necessarily very thin.

chemical means. The first process for the anodic exidation of aluminium was patented in 1924 by Bengough and Stuart, under the the advantage that it can be applied in min to a Department of Scientific and Industrial surface already blackened by exposure. Research. In this process the metal is made the anode in a 3% solution of chromic acid at about apprit rather than the letter by the development 40°C., using a neutral cathode, usually of stainless atecl; it is largely used for the protection of durelumin in aeronoutical work (Sutton and Sidery, J. Inst. Metals, 1927, 28, 24). Recent developments have been described by R. W. Buzzerd (J. Res. Nat. Bur. Stand. 1937, 18, 251). The process can be applied to other aluminimm alloys containing not more than 4% copper, and the alumina coating (which contains incorporated

well as corresion. It also has the property of acting as a mordant for dyes, and articles to treated may receive coloured finishes by immer sion in aultable dye solutions; alternatively the protection may be reinforced by lanoline or varnish, for which the coating provides an efficient hond. An advantage of the enodic as compared with cathodic methods of protection is that whereas large end ineccessible areas can readily be made anodic, there is great difficulty. as already pointed out (p. 387), in rendering a large aree effectively cathodic. More recently other anodic processee have been introduced with the more particular object of mereasing the intensity of the colours for decorative purposes, but this object is achieved at the expense of somewhat reduced efficiency on account of the increased porcenty of the coeting. These processes depend on the use of two main alternative electrolytes, solutions of sulphure end exalic acids respectively ("Alumilite" and Elexyl processes). Duplex baths containing chromic acid are also used; other patented processes employ small amounts of morganic salts for the production of coloured or black coatings its presence encourages further The treetment is followed by application of he underlying metal. "sealing compounds" for the purpose of fixing the colour and increasing resistance to corresion.

Corresion products other than oxide films are not often sufficiently protective to warmnt their etimulation by artificial methods. The green patina on copper is exceptional, and here the desire for the rapid production of the patina arises also from its asthetic value and the long period of some 10 or 20 years normally required for its development. An interesting artificial method (Vernon, J. Inst. Metals, 1932 153; ass also Freemen and Kirhy, Metals and Alloys, 1932, 3, 190) is that in which the specimen is made the anode in an electrolytic hath kept at 95°C. and containing magnesium sulphate and hydroxide, with potassium brom-ate as oxidising agent. In ac short a time as In minutes (or even sooner with appropriately aumiliar manner to that of the natural patina The industriel importance of aluminium and (supre). A further method (Vernon and its alloys led to various attempts to increase the Strond, B.P., 1938, under Department of thickness of the natural oxide film by electric Scientific and Industrial Research), consisting of spraying the aurface with thionyl chloride or other chlorine derivative of sulphuric acid, has

More commonly, neture is imitated in the of a reaction product which, whilst protecting the underlying metal, finds no counterpart in any natural process. Such, for example, is the case with the verious phosphale coatings on iron and atecl. These originated in 1906 in the process of Coslett (Birmingham), whereby the articles were treated in a hot solution of phosphone acid to which iron filings were added "Perkerising," employing a solution of iron and decorative (blackish) coating having many practical applications. "Bonderising" (much used in the automobile industry) is similar, hut the bath contains substances that greatly speed up the reaction and the coating is used mainly for increasing the adhesion and permanence of a subsequently applied paint or varnish.

The protection of magnesium and magnesium alloys, of importance in aeronautical construction, is limited almost entirely to chemical methods, with or without the subsequent appli-cation of paint. The "I.G. chromate dip" employs a hot solution of sodium dichromate in nitric acid, producing a yellowish iridescent film, the protective value of which is limited. A development of this is the alum-dichromate process of Sutton and Le Brocq (J. Inst. Metals, 1931, 46, 53; 1935, 57, 199). The selenising process of Bengough and Whithy (bid. 1932, 48, 147; Trans. Inst. Chem. Eng. 1933, 11, 176; D.S.I.R. Rep. Chem. Res. Board, 1934-7, London, H.M.S.O., 1938) involves immersion of the metal for a few minutes in a solution of selenious acid or acidified sodium selenite and produces a surface film of selenium which has considerable self-healing properties. Under test with sea-water spray the difference in behaviour hetween coated and blank specimens is spectacular, for the latter may disappear entirely whilst treated specimens remain comparatively unaffected. In hoth the alum-dichromate and the selenium processes (hoth of which are covered by British patents) the coating appears to find its most promising application as a hase for paints.

Reference (Natural and Artificial Films) .-E. S. Hedges, "Protective Films on Metals," 2nd ed., Chapman and Hall, London, 1937.

Paint and Allied Coatings.—The preservation of iron and steel hy means of paint, for which, according to Jordan and Whitby (infra) some 40 million gallons of paint are used annually in Great Britain alone, involves many factors associated hoth with the paint and with the metal. A paint system is complex and its components-pigment, vehicle, thinner and drier-are interdependent in their contributions to the properties of the whole. It is usual, however, to classify protective paints according to the effect which the pigment may be expected to have per se on the corrosion of iron. Three types may he recognised, according as corrosion is likely to be (I) inhibited, (2) unaffected, (3) stimulated, hy the pigment alone.

Characteristic examples of inhibitory pigments are provided hy chromates (e.g. basic lead chromate, zinc chromate) and in this case the mechanism almost certainly resembles that of similar inhibitors in immersed corrosion, tending to render the iron passive. They are most usefully employed as "priming paints," in immediate contact with the metal. By far the most widely used primer is red lead, but here the passivating effect is probably supplemented by the forma-tion, by reaction with the linseed oil, of lead compounds of high molecular weight which serve to cement the particles of pigment have the advantage, to an even more marked together. In practice, litharge normally present extent than graphite, of a "flaky" or "leafy"

and manganese phosphates, gives a protective this reaction. "Non-setting" red lead, having the minimum content of litharge, has been deprecated for corrosion-protection, particularly of ships' hulls (A. McCance, Trans. Inst. Eng. Scotland, 1936, 333); on the other hand, J. C. Hudson states (Proc. Chem. Eng. Group, S.C.I. 1937, 19, 32) that satisfactory results have been obtained from non-setting red lead in the Iron and Steel Institute Corrosion Committee's tests. Probably the desideratum is a compromise in which the optimum proportion of inhibitory and setting capacity (depending on the type of work to be protected) shall he retained. For many purposes red lead may he safely diluted by the addition of varying proportions of inert "extenders," e.g. barytes, or asbestine.

Priming paints usually need to be reinforced hy a top coating either for the purpose of healing mechanical defects in the undercoat or by effecting a more efficient exclusion of the surrounding liquid or atmosphere. For this purpose pigments of either groups (2) or (3) may be used. Red iron oxide is used extensively and confers good durability. Intrinsically neutral in character it may function either as an inhibitor or stimulator according as it is associated with alkaline or acid impurities from its manufacture. Recent developments include the use of red iron oxide with red lead or zinc chromate (mixed pigment) to give a combined priming and finishing coat. White lead (basic lead carbonate) and zinc exide have mild inhibitive properties, but are more appropriately included with pigments of the second class; excellent as finishing coats for many purposes, they should not he used as primers where corrosion conditions are severe. The essentially chemical and physical mechanisms underlying protection hy red lead and red iron oxide have been shown hy lahoratory experiments (Lewis and Evans, J.S.C.I. 1934, 53, 29) and hy field tests (Britton and Evans, J.S.C.I. 1932, 41, 215); they have also heen demonstrated hy an electrochemical method (time-potential curves) hy Burns and Haring (Trans. Electrochem. Soc. 1936, 69, 169).

Pigments of the third class include such substances as graphite and certain metallic (e.g. bronze) pigments, which, on account of their cathodic nature, cannot normally he employed in direct contact with the metal. Their value lies either in their good weathering properties or in their ability to exclude moisture, and hence they are employed with advantage as finishing coats. In the case of graphite, exclusion of moisture is due largely to the 'flakey'' nature of the particles, the flakes lying on the surface with their long axes parallel to the metal. This property is also possessed by micaceous iron ore, a neutral pigment that is stated to he specially resistant to tropical conditions. Aluminium in the metallic condition might he expected to undergo "sacrificial corrosion" in contact with iron, but in practice the particles are probably sufficiently oxidised to render them neutral in character; they also in commercial red lead is mainly responsible for nature. Metallic lead pigments are relatively

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ditious. Of available media, linseed oil is by far the most widely used. Heat-treatment of the oil reduces the permeability of the paint film to water vapour and also its liability under certain conditions to act as a depolariser for pascent bydrogen and hence to facilitate corresion. Moreover, the alkali reproduced at eathodes during corresion in salt solutions is more likely to produce "alkaline peeling" of the paint film if raw, and hence more saponifiable oil is employed. Permeability to water vapour may be atill further reduced by the addition of certain gums, waxes or resins. For special purposes, where high permeability to water is required, tung oil may he wholly or partially substituted for linseed oil. For most oil paints there is probably an optimum ratio between medium and pigment content (J. N. Friend, J. Oil Col. Chem. Assoc. 1922, 5, 276), hut if the pigment is inhibitive (e.g. red lead) rather than non-unhibitive (e.g. red iron oxide) lower proportions are permissible without detriment to corrosionprotection (Britton and Evans, J S.C I. 1932, 51, 211T). Jordan and Whitby (infra) recommend, however, even with inhibitive primers, a pig ment concentration as high as is consistent with penetration of the paint into all crannies and fregularities. The proportion of "thinner" (if this is the usual turpentine or alternatively white spirit) is important from the corrosion protection stand point prohably only in so far as it controls the thickness of the resulting paint film. "Drying" of the linseed oil in a paint film is catalyeed by the "drier" (certain compounds of lead, cobalt, or manganese) of which no more should he used than is necessary to ensure the drying of the paint within a reasonable time, depending on the conditions of axposure. A greater proportion of carefully selacted driers as usually employed in paints for under-water service (see under "Corrosion of Metals—Corrosion of Ships" Hulls," supra).

The preparation of the metal surface previous to painting is of greatest importance. If the paint coating is to have maximum life, all millscale or rust should be removed. The danger of mill scale is its liability to peel on exposure carrying with it en bloc the overlying paint. As, however, the scale is itself intrinsically resistant, there is hope of progress in the production of a thin scale that will have no tendency to peel. The danger of rust hes in its hygroscopic and adsorptive character. If electrolytes are present on the anriace beneath the paint film this may act as a membrane through which mosture may pass osmotically from outside, even though the paint may be thoroughly dried hefore exposure (Bartell and Van Loo, Ind. Eng. Chem. 1925, 17, 1052). Removal of scale may be effected (1) naturally, by weathering, (2) mechanically, by cand blasting (wire brushing is more austable for the removal of rust), (3) chemically, by picking processes. Insufficient weathering often causes trouble, since the flaking of residual ecale may continue after the paint is applied. Sand blast-

¹ For the chemistry of the drying process and of paints generally articles on "Paint" should be con-suited.

inert and are highly durable under most con-ling has certain limitations, but has the obvious advantage of rapidity. J. C. Hudson (Vth Rep. Corr. Ctte. Iron and Steel Inst. 1938) states that pickling, as compared with partial weather. ing and careful hand-cleaning, increased the his of paint films by more than 150%. Pickling is usually carried out in a hot dilute solution of either hydrochloric or aulphuric acid, preferably the latter, with the addition of an inhibitor for the purpose of confining the attack to the scale and protecting the metal itself. Adequate wash ing, preferably in water containing alkal, chould follow picking. The use of phosphone acid, at suitable dilution and temperature, has been advocated on the grounds that it leaves a film of basic iron phosphate, which increases the adhesion of paints, and washing may then be dispensed with. Alternatively, scale may be removed by the cheaper sulphune seid, and s amalier amount of phosphoric acid applied after an intermediate washing (H. B. Footner, Vth Rep. Corr. Ctte. ibid. 369). In any case the aurface should be thoroughly dry before painting is commenced. Moreover, adequate time should be allowed for any coat of pant to "dry" or set before the succeeding coat is applied. The foregoing remarks have referred particularly to iron and steel surfaces. The painting of non forrous metals may present epecial considerations, usually in the direction of providing a suitable chemical hond to ensure the adhesion of the paint; certain available methods have been discussed above. Trouble is frequently experienced in the painting of galvanued surfaces (hot-dipped zine coatings); this may be overcome either by a preliminary weathering of the sheets, or by a suitable atching treatment (Vth Rep. Corr. Cite, ibid. 303); see also "Hot dipping Processes", p. 306. Grease Coatings and Slushing Compounds -For the temporary protection of iron and steel articles, machine parts, etc., during fabrication or storage, coatings of grease are employed which may subsequently be removed by an organic colvent. For this purpose landing is especially efficaceous and is usually employed in the form of solution in white spirit or solvent naphtha, the evaporation of which leaves the grease film uniformly distributed. Alternatively the grease may be applied in the form of an emulsion with an aqueous colution of acdum dichramate, or even with soap solution, such

mixtures being generally known as "slushing compounds." References (Paint Coatings).—Jordan and Whitby, "The Presexvation of Iron and Steel by Means of Paint," 1936, Research Association of British Paint, Colour, and Varnish Manufacturers, Sixteenth Bulletin; L. A Jordan, The Preparation of Metal Surfaces for Paint rng," Chem. and Ind 1937, 56, 361. (For Protection of Ships' Hulls, including Anti-Touring Preparations, see Cornosion, General,

p. 383, 383). Lacquers and Varnishes .- For protection against tarnishing of polished aurfaces, eg. of copper and alver and their alloys, transparent films of lacquer or varnish are employed. Formerly a natural ream (usually aheliac) was the invariable component; fn a "lacquer"

methylated spirit) the rapid evaporation of which yielded a film dry to the touch very quickly after application; in a "varnish" the medium was a drying oil (incorporated by heattreatment) the setting of which occupied a much longer time, the resulting film, however, being tougher and more elastic. Whilst this distinction between a lacquer and a varnish is still roughly true, natural resins are now very largely supplemented by a wide range of so-called synthetic resins. (For detailed information, see "Synthetic Resins and Allied Plastics," by Barry, Britton, Langton, and Morrell, London, Oxford University Press, 1937. Also "Modern Developments in Plastics," by Sir Gilbert Morgan, Chemistry and Industry, 1937, 56, 103). These were at first of phenol formaldehyde type, available only in the form of the simple spirit solution (e.g. "Novolak") but many types of oil-soluble resins are now employed in the manufacture of protective varnishes. of the modified phenolic derivatives find use in protecting metal surfaces against industrial fumes. They are also used as stoving lacquers for the protection of tinplate against acid juices, their freedom from odour and taste rendering them particularly valuable for the purpose. A. F. Brockington (J. Electrodepositors' Tech. Soc. 1937, 12, 133) states, however, that the modified alkyd types of phenol aldehyde resins are seldom used in the unpigmented state on metals since they have an acid reaction and tend to tarnish copper and brass during stoving. Other oil varnishes employing synthetic resins (e.g. those obtained from polystyrene and coumarone) have good alkali-resisting properties.

"Cellulose lacquers" consist usually of a solution of cellulose nitrate in a mixture of solvents (amyl acetate, etc.), but in modern practice either natural or synthetic resin is usually incorporated for the purpose of improving the adherence and gloss of the dried film. The mixture also includes, in common with other "cold lacquers," a certain proportion of higher alcohols, their function being to prevent too rapid volatilisation of the solvent with consequent "blooming" or "blushing" of the film; this term refers to a defect caused by precipitation of moisture when a lacquer is applied under humid conditions. Cellulose lacquers, commonly applied by spraying, are particularly suited for highly polished surfaces. Both cellulose and gum laoquers may be tinted with suitable dyes. The varnish-like nature of certain invisible oxide films (p. 392) may be recalled; it has been suggested (Vernon, Res. Rep. Brit. Non-ferrous Met. Res. Ass., Nov. 1927, 9) that the production of such films by heat-treatment (e.g. on brass) might with advantage precede the application of lacquer, thus reinforcing the protective effect (partly by providing a better chemical "bond") whilst retaining the natural colour of the original surface. The application of lacquers and varnishes to thicker (visible) films produced by chemical treatment is common practice; they may either supplement the protective effect of

this was dissolved in a volatile solvent (e.g., and magnesium alloys, p. 392) or they may serve to protect a decorative coating (as produced in certain metal-colouring processes) which itself may have little protective value.

The lacquering or varnishing of tinplate for food containers has for its object not so much the supplementing of the protection of iron by tin as the suppression of certain reactions between tin and fruit juices which may cause either blackening of the tin or aiscoloration of the juice. Indirectly, indeed, the lacquer coating, if imperfect, may lead to local attack on the tin, and by concentrating the attack at the weak places may bring about the perforation of the container. Walker and Lewis (J. Ind. Eng. Chem. 1909, 1, 754) suggested that in such cases a varnish film might, by acting as a hydrogen acceptor, facilitate the depolarisation of hydrogen even in the absence of oxygen and so accentuate the anodic attack on the adjacent bare places. Morris and Bryan (D.S.I.R. Food Investig. Reps. 1933, 145) conclude that the lacquer film in modern practice is inert although the tin coating beneath may play some part electrochemically. They recommend (ibid. 1935, 169) supplementing the usual pre-fabrication lacquering of the tinplate with a final lacquering of the can interior with a quick-stoving lacquer.

Chlorinated rubber has within recent years received considerable attention as a protective coating (G. Schultze, Korr. u. Metallschutz, 1936, 12, 249; J. P. Baxter, Chem. and Ind. 1936, 55, 407). In the form in which it is typically applied—a solution in xylene or other solvent with incorporated plasticisers such as certain chlorinated hydrocarbons—it is analogous to a synthetic resin lacquer. Pigments may, however, be added, with or without a proportion of drying oil (e.g. tung oil) and hence certain mixtures resemble oil paints. essential feature of the chlorinated rubber coating, besides its high elasticity, is its capacity of giving a high resistance to chemical attack, both acid and alkali; it appears to be well suited for under-water protection, particularly

under tepid conditions, as in thermostats, etc. Bituminous Coatings .- The simple "bituminous paints" are not true paints as previously defined. They consist of either natural asphalt or pitch (petroleum residual or coal tar) dissolved in a volatile spirit (e.g. white spirit, solvent naphtha, etc.) so that drying, as distinct from the "drying" of an oil paint, consists merely in the evaporation of the solvent. Such bituminous coatings afford good protection to iron and steel, sometimes under conditions such as would be deleterious to ordinary oil paints (e.g. atmospheres polluted by chemical fumes); they are, however, adversely affected by excessive sunlight. They are widely used for structural work, for which purpose they are improved both aesthetically and in durability by pigmentation, particularly with aluminium powder. Excellent results, for example, have been quoted for the use of tar paints containing 12 to 15% aluminium (Vth Rep. of Corr. Ctte., Iron and Steel Inst. 1938, 320); the aluminium floats to the surface of the film which, when dry, has practically the same appearance as an aluminium paint the primary coating (e.g. protection of aluminium | proper. A modification of bituminous paint,

using bitumen together with drying oil additions, | mediate layers can be detected, each character. using nitimen together with drying oil additions, is medieto layers can be detected, each character her coating being "atord" after appheaton, ised by an intermetallic compound. Advantage results in black "iapan." For the protection is claimed for a process of subsequent annealing of pipelines underground, cool-tar pitch, the bass of the well known "Angus Smith compound," is extensively employed. The original pound, "is extensively employed. The original patent (1818) appended the use of inneed oil as exact the control of the an addition to the pitch to obtain the right consistency and elasticity. In modern prectice the pipes are heated end immersed in the molten hath of refined coal tar pitch (sometimes with additions, for example, are not normally made certain "filler" additions) end frequently the coating is reinforced with bessian or similar quantities of either aluminium or tin are meterial. Demage to this protective coating in treasit from the mill to the trench may result in local failures in aervice. For marine con-ditions, J. N. Friend (D.S.I.R. "Deterioretion application in the menufacture of food containers of Structures in Sea-water," XVth Rep. Ctte. Inst. Civ. Eng. 1935, 85) reports excellent results from the use of coal ter coatings on iron and ateel, end states that coal-tar from horizontal retorts was superior to that from vertical retorts, whether applied hot or cold, it was improved by the addition of alaked hme (see also "Corrosion and Protection of Ships' Hulls," p. 382). "Bituminous Materials in Paint" have been eurveyed by L. A Jordan (Res. Assoc. Brit. Paint Manufac. 14th Bull. 1935); see also Jordan and Whitby, 1936, supra.

Metal Coatings .- The non-ferrous metals most commonly applied in the form of thin coatings for the protection of iron and attel are aluminum, lead, tm, and zinc Methods of appheation include either purely metallurgical processes (rolling, hot-dipping, cementation, arraying) or electro deposition.

"Rolled-on" Contings — In the "old Sheffield

plate" process (rendered obsolete by the intro duction of adver plate) thin sheets of adver were duction of salver plate) than sheets of silver were rule that applies to all metaline coatings, wance beaten on to copper, the operation being frequently depend for their corrosion resistant assisted by host. Within recent years the principle silver reviewed in "metal clading" processes. Thus, copper-clad and notekt-clad patcels are produced by rolling or drawing actels are produced by rolling or drawing a steels are produced by rolling or drawing a heated builds of atcel. "and without the cating itself is contributing heated builds of atcel." Aluminum clad the formation of the protective layer, and its initial thickness should thus be at least sudicions. ateel has been proposed as a aubstitute for tinplate in the conning of foodstuffs. A promising development is the rolling of a veneer of atamless conditions which result in "ascrificial" corresion. steel on to an ordinary steel base Aluminium and hence loss of the protective coating, should alloys of the duralumin type (having high strength but a susceptibility to intergranular corrosion from which the weaker aluminium is free) are similarly covered with pure aluminium. The duplex material (American "Alclad," "Aldural," German "Allautal") is British used extensively in aircraft. Alternatively (as in "duralplat"), a relatively resistant airconeluminium alloy (free from copper) may be freshly used for the outer layer, the rather less resistance p. 391). to corrosion being compensated by the greater resistance to weer.

Hot Dipping Processes -Hot dipping is em. How dryping if the property of underlying metal is provided by a certain galvanised iron, but pointing is usually neces-amount of alloying. Exceedingly thin on tin, sary: this is facilitated by the readiness with the alloy layer reaches much grosser dimensions which paint adheres to the material without any on zinc, where two and occasionally three inter. prehminary treatment.

set up by the presence of these layers. In all cases scale adhering to the metal must be completely removed by picking prior to the hot-dipping. The composition of both basis and covering metal is usually more rigidly controlled for timplate than for galvenused iron. Metallic to the tin bath, but to the zine bath very small commonly made with the object of increasing the fluidity of the bath or improving the appear and dairy utensils, whilst gelvanised fron is used for roofing and for much large-scale work exposed to the open air. (Hot tinning is also applied to copper articles, porticularly water pipes and cooking utensils.) Tim and zinc ere normally cathodic and anodic respectively to iron (see " Electro-potential Series," Table III, p. 376). Hence, in the event of the underlying metal becoming exposed at any point, in the presence of a austable electrolyte, the iron in contact with tin is attacked preferentially, leading to perforation of the sheet, whilst a zino coeting, under similar conditions, will undergo "sacrificial corrosion" where the underlying metal is exposed and thus protect it from attack. Circum atances sometimes arise, bowever, in which a reversal of polerity takes place, as, for example, with timplate in the presence of certain fruit mices. It is important that a certain minimum thickness of coating should be exceeded if a reasonable life is to be assured. This is a general rule that applies to all metallic coatings, which frequently depend for their corrosion resistance to aurvive this interim period. This applies equally to coatings of the anodic type, since the be avoided.

Galvanised iron, when used for structural work exposed to the open air, is frequently further protected by painting. The unsatis factory adherence of paint to the galvanised coating may be overcome by a preliminary period of weathering; various solutions (eg oxalic acid), are also employed for etching the freshly deposited coating prior to painting (see

Terne plate consists of iron or steel sheet costed (hy hot-dipping) with a lead alloy containing 10 to 25% tin. (Leed itself does not give a satis-

Cementation Processes.—The principal cementation process (heating of the previously cleaned basis metal with the coating metal, the latter in powdered form, usually "diluted" with oxide) is that of "sherardising," due to the late Sherard Cowper-Coles, in which iron or steel is coated with zinc. The process is particularly suited for small articles, such as bolts, nuts, screws, for which it has the advantage over hotdipped coatings in respect to a considerably smaller dimensional change. The coating differs principally from the hot-dip coating in that there is usually only one alloy layer, whilst the outer layer of pure zinc is often not represented at all. It is in the uniformity and continuity of the alloy layer that the corrosion resistance of sherardised coatings principally This alloy is normally anodic to iron but less so than zinc.

The analogous process of cementation by means of aluminium (calorising) is used more particularly for protection against oxidation at high temperatures, where the protection is due to the refractory properties of the film of alumina. The coating is also useful for protection against

sulphurous fumes.

Metal-spray Processes.—The process devised by Schoop in 1910, whereby a spray of finely divided metallic particles is ejected from a blowpipe or "pistol," has within recent years assumed important industrial dimensions. Metal wire is supplied to the pistol, in which it is melted at the tip by an oxyacetylene flame, atomised by excess of gas, and carried forward at a relatively low temperature (actually below the ignition point of a match). Adhesive deposits of various metals may be sprayed on to steel, which should be previously sand-blasted to obtain the necessary "keying," since no appreciable alloying occurs. An obvious advantage of the method is its ability to apply a protective metal coating to structures, in situ. Recent modifications, as the Schori processes, have aimed at increasing the speed of working by supplying metal powder to the pistol instead of wire; this process also sprays metal/glass mixtures which are claimed to provide flexible coatings of metal silicate resistant to sea-water attack. Marked resistance to corrosion in industrial and marine atmospheres has been obtained by the spraying of thin coatings of aluminium on steel by the ordinary method, particularly when the coatings have been subsequently treated with cellulose lacquer (W. E. Ballard, Trans. Manch. Assoc. Eng. 1935, 113). The possibility of using sprayed metal coatings for the protection of ships' hulls is referred to in the Fifth Report to the Iron and Steel Institute Corrosion Committee (l.c., 219).

Electro-deposited Coatings.—Much research has been carried out with the view to increasing the efficiency of electro-deposition processes, by which a wide range of protective metal coatings are now produced. It has been shown that interalloying of coating and base does not take place, although in certain cases slow diffusion may occur subsequent to deposition. On the other hand, the relationship may be so intimate that the lattice of the basis metal may

by the deposit (A. W. Hothersall, Trans.

Faraday Soc. 1935, 31, 1242).

Electro-deposition has the advantage of producing a uniform coating under controlled conditions with little dimensional change. It is particularly suitable for small articles of irregular shape, but for special purposes may be applied to large-scale work. Small articles are frequently plated in bulk by the "barrel" method; a rotating barrel dipping into the electrolyte contains the articles, the mutual contact of which ensures electrical connection and provides a certain burnishing action which improves the appearance of the deposit.

Zinc coatings are produced commercially by

each of the four main methods; the term "galvanised" iron, however, almost invariably refers to the "hot-dipped" coating. Tin coatings are produced to a comparatively small extent by electro-deposition, which has, however, been recommended for supplementing the hotdip process, the two operations yielding a coating

that is practically free from pores.

Nickel-plating on iron and steel is carried out the largest scale. Excellent protection against corrosion is afforded under many conditions, nickel coatings being particularly suited for parts of equipment which must resist both corrosion and abrasion. In atmospheres containing traces of sulphur dioxide, provided a critical relative humidity of approximately 70% is exceeded, the polished surface undergoes a characteristic filming or "fogging" (see General article under Atmospheric Corrosion). In modern practice nickel coatings are covered, for decorative purposes, with a film of electro-deposited chromium. This resembles nickel in its extreme hardness and resistance to wear, fine appearance, and ability to take a high polish, but has the advantage of freedom from fogging. (Small amounts of chromium alloyed with nickel also inhibit fogging, but the two metals cannot, as yet, be deposited simultaneously.) An analogous recent development is the plating of a thin film of rhodium on electro-deposited silver. This retains the advantages of silver plate with freedom from tarnishing, to which unprotected silver is characteristically liable in atmospheres containing traces of sulphur compounds.

The use of electro-deposited cadmium is increasing considerably. Whilst it appears to have no appreciable advantage over zinc for open-air conditions, its whiter and pleasanter appearance, particularly when polished, is advantageous for indoor purposes, as in electrical and radio parts. In the Udylite process, iron articles, after electroplating with cadmium, are heated at 150° to 200° for several hours.

Arising from difficulties in producing coatings of lead by other methods (e.g. absence of alloying in hot-dip and cementation processes and health hazards in spraying processes) developments have recently taken place in the large-scale electro-deposition of lead on iron for roofing and constructional purposes (F. W. Hay, Metallurgist, 1937, 11, 23). High resistance to industrial atmospheres is brought about by the formation of a surface film of insoluble lead sulphate. be continued and the structure exactly copied Developments in electroplating are surveyed by

S. Wernick (Chem. and Ind. 1934, 53, 948; belong to the rhombohedral system, and are Ind. Chem. 1935, 11, 233). See also works isomorphous with hæmatite (Fe O3). They recommended below.

TESTING OF METALLIC COATINGS .- Many methods are in use for testing soundness, thickness, and adhesion of metal coatings. One of the earliest schemes for detecting pinheles was by means of the ferroxyl indicator of Cushman and Walker. This is prepared by the addition of a little potassium ferricyanids and phenolphthalein to a dilute solution of sulphuric acid; a convenient method of carrying out the test is to use a test method of carrying out the test's to use a test, paper saturated with the reagent, which is pressed on the surface. In the case of a cathodic coating (e.g., nickel nn iron) the position of the punholes is inducated by a blue spect of services ferrigrands where nron passes into solition. For timplate, Macanaghian, Clarke, and Prytherth (d. Iron and Steel Inst. 1022, 123, 159) recommend a solution containing sodium chloride with potassinm ferricysmide; they also obtain excellent results from a simple "bot water test," observing suitable precautions, Pinholes in anodio coatings (e.g. 2100 on 110n) are more difficult to demonstrate, but usually they are of less consequence in practice. The thickness of zino deposits on iron is still determined by the old established Preece test; this consists in repeated immersion of the samples (usually galvanised wire) in copper sulphate solntion at one minute intervals, the number of immersions required to produce a deposit of copper on the basis metal giving an indication of the thickness of the costing. A number of methods have been proposed for overcoming certain objections to the Preece test. Britton (J. Inst. Metals, 1936, 58, 211) makes the sample the snods of an electrolytic cell at a fixed current density and notes the time taken to remove the costing from a given area. The thickness of tin coets on iron is determined rapidly by Clarke by removal in a colution of hydrochlorio acid and antimony chloride, the latter preventing attack on the iron. In the "B.N.F. jet test" the thickness of the coating at any desired point is determined by measuring the time required for perforation of the coating by a jet of an appropriate solution which is allowed to impinge on the surface at a constant pressure. It is applicable to a wide range of coatings (S. G. Clarke, J. Electrodep. Tech. Soc. 1936, 12, 157; 1937, 12, 1-18). Methods of testing zinc coatings have been comprehensively surveyed by L. Kenworthy (J. Inst. Metals, 1937, 61, 143) References (Books dealing with Metal Coatings)

References (Books dealing with Metal Losungs),
H. S. Rawdon, "Protective Metalthe Coatings,"
Chemical Catalog Company, New York, 1928,
H. Bahlk, "Galvaniang, 2ad ed. (Epglah
translation), E. & F. N. Spon, London, 1836;
E. S. Hedges, "Protective Flints on Metals,"
2nd ed., Chapman and Hall, London, 1037.
W. H. J. W.

chloride.

ABRASIVES and CORUNDUM.)

vary considerably in habit, having the form of hexagonal prisms, pyramids, or tables. In some crystals the combination of six faces of the primary rhombohedron with two faces of the basal pinacoid produces a form strikingly similar to the regular octahedron. There is no true eleavage, but often the crystals and crystal. line masses show a pseudo-cleavage or pering parallel to the basal plane or to one or more faces of the primary rhombohedron, due to lamellar twinning in these planes. On these aurfaces the lustre is sometimes pearly, but ntherwise it is usually vitreous, or inclining to adamantine in character. The fracture is un even to conchoidal; and the sharp jagged edges of the irregular groins give effect to the cutting power of the material when used as an citizing power in the material when used as an abrading agent. Analyses show the presence of 95-99% of alumina, with small amounts of fermo oxide, magnesia, silca, water, etc, and sometimes traces of ehronium. Sp gr. 39-41. The crystals are optically uniaxial and negative; refractive indices for sodium light \$\omega\$1.768, \$\epsilon\$1.760. The dichroism is marked. Next to diamond nt is the hardest of minerals, heing placed as No 9 m the scale of hardness , on this account it was formerly known as adamantine spar,

A distinction is made between the transparent precious corundum" and the dull, oreque "common corundum," while a third still less pure variety is known as emery (v. ABRASIVES). This division slso corresponds with the economic uses of the mineral, "precious corundum" being used for gem stones, and "common corundum " and emery for shrasive purposes The pure, well crystallised material is perfectly colourless and transparent with a high degree of hrilliancy. Traces of morganic impurities, however, produce a wide range of colours, which are more vivid in the stones of gemquelity, namely red (ruby), orange and yellow ("oriental topsz"), green ("oriental emerald"), hlue and indigo (sapphire), and violet ("oriental emethyst"). Although such names are m popular use for tewellery, this mineral has, of course, no relation to topaz, etc. Parti-coloured atonca are not uncommon. The colour of

common corundum is usually grey or brown. Corundum is not a widely distributed mineral, but it is found in ahundance at certain localities It occurs in igneous and metamorphic rocks in which no free ailica is present as quartz, such as syemite, gness, peridotite, and serpentine, and in crystalline limestones. Material of genquality is usually collected as water-worn pehbles in gravels derived from these rocks Material for gem-stones and that for abrasics purposes are net, as a rule, mined in the same localities. Taking first the localities for the gemvarieties, the following may be noted as a few of the most important. The best rubies are CORROSIVE SUBLIMATE. Mercuric obtained from the famous mines near Mogok corubin. An artificial corundum (v. erystelline marble near its contact with guess; they are mined in the detrital deposits derived APRASYES and CONVACAN, we will be a seen and the seek Dark-coloured rules are from the Hindl kurund, Sanskrit kurarindo, bottamed with aspphire from alluvial deposit Crystallized slumma (Al-Q.). The crystals in the provinces of Chantabus and Krat is Macon Co., North Carolina. The best sapphires come from the gem-gravels of Ceylon; other localities of importance are Siam and Kashmir. Dark blue and greenish-blue sapphires occur abundantly in gravels near Anakie in Queensland; while paler stones of various shades "fancy stones") are plentiful in the neighbourhood of Helena in Montana.

Common corundum as used for abrasive purposes (apart from emery) was formerly mainly mincd in the United States; later the Canadian and South African deposits came into prominence; while now the latter are being ousted by the artificial products carborundum and alundum. India has also since remote times produced a certain amount which has been largely used by the native lapidaries. In the United States the corundum deposits extend along the Appalachian Range from Alabama to Massachusetts, reaching their greatest development in North Carolina and Georgia. Here they occur in peridotites near their contact with gneiss. In Canuda a belt of corundum-bearing syenites (nepheline- and related alkali-syenites); associated with the Laurentian gneisses, extends for 100 miles across Renfrew, Hastings and Haliburton across Renfrew, Hastings and Haliburton counties in Ontario, the principal mining district being near Craigmont in Renfrew Co. Large rough crystals of corundum are abundant in gneiss in Madagascar and at Steinkopf in Namaqualand, South Africa. More recently, important deposits have been discovered and worked in the Zoutpansberg and Pietersburg districts in northern Transvaal.

The corundum-bearing rock as mined is crushed between hardened steel rolls and the heavier corundum separated in water by means of jigs and concentrating tables. It is sifted into different grades of fineness and used (1) as loose grains; (2) mixed with glue or other soft coment as a coating on paper or cloth; and (3) as corundum-wheels for grinding. The last-named are of three classes: (a) The "vitrified wheel," in which the corundum is mixed with clay and baked at a high temperature, with the result that the grains of corundum are set in a matrix of porcelain. (b) The "chemical wheel," in which sodium silicate (c) The "cement is the binding material. wheel" with shellac, rubber, linsecd-oil, etc., or magnesium oxychloride. Another use for corundum, depending on its hardness, is for the pivot supports for watches (watch-jewels) and other delicate instruments, the poorer qualities of gem-material being used for this purpose. Although corundum is the richest ore of aluminium, there are difficulties in its use for the extraction of the metal. It has, however, been used directly in the preparation of aluminium-

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copper and aluminium-iron alloys.

The artificial preparation of crystallised corundum was first effected by M. A. Gaudin, in 1837, by decomposing potash-alum with charcoal. It has since then been produced by a variety of methods, and is now manufactured

Siam. Stones of good quality have also been ruby were prepared by E. Frémy and Feil, in mined together with common corundum in 1877, by heating a mixture of alumina and redlead with potassium dichromate in a porous fireclay crucible; a fusible aluminate of lead was thereby formed, which was decomposed by the silica of the crucible with the separation of crystallised alumina. The crystals so obtained, though of some size, good colour, and perfectly transparent, were unfortunately too thin for cutting as gems. A far greater measure of success was achieved by A. Verneuil in 1902, who has produced fine gems of all colours. Interesting experiments from a petrological point of view were conducted by J. Morozewicz in 1898, in which he obtained crystallised corundum in silicate magmas supersaturated with alumina. Artificial corundum, used as an abrasive under the trade name of alundum (aloxite, adamite), is now manufactured on a large scale at Niagara Falls by fusing bauxite in an electric furnace. Corubin formed as a by-product in the Goldschmidt thermite process

is also an artificial corundum.

References.—A. E. Barlow, "Corundum, its Occurrence, Distribution, Exploitation, and Uses," Canada, Dept. Mines, Geol. Survey, Mem. 57, Ottawa, 1915, 378 pp. (with special reference to the Canadian deposits); T. H. Holland, "Corundum," Manual of the Geology of India, 2nd ed., Calcutta, 1898 (with special reference to the Indian deposits); A. L. Hall, "Corundum in the northern and eastern Transvaal," Mem. Geol. Surv. South Africa, 1920, No. 15; G. P. Merrill, "The Non-metallic Minerals," 2nd ed., Now York, 1910; J. H. Pratt, Bull. U.S. Geol. Survey, 1906, No. 269. J. H. Pratt, 2nd ed., Now York, 1910; J. H. Fratt, Bull. U.S. Geol. Survey, 1906, No. 269; J. H. Pratt and J. V. Lewis, North Carolina Geol. Survey, 1905, i.; M. Bauer, "Edelsteinkunde," 3rd ed., by K. Schlossmacher, Leipzig, 1928–1932, and English translation, "Precious Stones," by L. J. Spencer, London, 1904; "Abrasives," Imperial Institute, London, 1929.

L. J. S. CORYBULBINE v. Corydalis Alkalords. CORYDALINE SUB-GROUP.

CORYCAVINE v. Corydalis Alkaloids, CORYCAVINE SUB-GROUP.

CORYDALINE v. Corydalis Alkaloids, CORYDALINE SUB-GROUP.

CORYDALIS ALKALOIDS .- The corydalis species belong to the Fam. Papaveracem: there are about ten different species known to contain alkaloids, of which C. cava (syn. tuberosa) is the most important (v. Wehmer, "Die Pflanzenstoffe," Jena, 1929, p. 388). For references to early chemical investigations, v. Ziegenbein (Arch. Pharm. 1896, 234 492). The most important alkaloid corydaline was isolated by Wackenroder as early as 1826.

Extraction .- According to Gadamer (Arch. Pharm. 1902, 240, 21) the finely ground corydalis roots are exhausted with 94% EtOH. The solvent is removed by distillation, the oily residue is acidified with acetic acid and diluted with H₂O. The aqueous solution is made alkaline with ammonia and extracted with Et₂O, when all alkaloids pass into the solvent except corytuberine, which is obtained by on a large scale both for use as gem-stones and evaporating the mother-liquor. By concen-for abrasive purposes. Beautiful crystals of trating the Et₂O-solution a first crystalline

fraction (a) melting from 160-180° is obtained which contains croydaline, bulbocapanine, corg-carine, and corybubine. This mirture may be exparated by successively extracting the cry-staline mass with smell emounts of boiling EtOH, The ethereal mother-liquor fcom (a) is then completely evaporated, when a fraction (b), melting from 130-140°, is obtained, which consists almost entirely of corydaline. The fraction (b) is converted into the bydrobromides, when first bulbocapnine HBr separates; by fractional precipitation of the remaining hydrobromides with ammonia corydaline, corybulbine, isocorybulbine, corydine, and coryeavine, and further emounts of hulbocapnine are obtained. The mother liquors are acidified with HGI when further crystalline fractions are obtained. The residue is transformed into the rhodomides. from which hy extraction with EtOH corycavemine rhodanide is left behind, Godamer's process is merely o general guidanco

and not an accurate prescription; it has often been modified if a special alkaloid was sought after. The separation and purification of the different alkaloids is a very tedious and laborious process; also the yields of the different alkaloids

vary to a great extent

Other investigated corydalis species are:
C. ambigua (Chinese C): Makoshi (Arch.
Phorm. 1998, 246, 381), Chou (A, 1928, 927;
ibid. 1929, 477, 1055; ibid. 1933, 652; ibid.
1934, 1014). Huang Munlon (Ber. 1936, 69,

[S], 1737) (important paper) C. ternys (Japanese C

Asaluna and Motigase (J.C.S. 1921, 120, I, 86). C. decumbens : Makoshi (Arch Pharm. 1908, 246, 401); Osada (Amer Chem Abstr. 1928,

22, 592) C. lutes and C. nobilis : Schmidt (Arch.

Pharm. 1908, 246, 575).

C. aurea: Heyl (Apoth Ztg. 1910, 25, 137); Eppson (Amer Chem. Abetr. 1935, 29, 2662) C. solida: Haars (Arch. Pharm. 1905, 243,

154). C. ternata Nakai (Korean G.). Go (Chem. Zentr. 1930, I. 234). The arrangement of the corydelis alkaloids

into three groups is due to Gadamer, Ziegenbein, and Wagner: Group I: consists of weally basic alkaloids.

which are readily exidised by iodine to berberinelike compounds (corydaline sub group).

Group II: includes atronger bases which are ettacked by iodine solution (corycavme aubgroup).

The alkaloids of Group 11 ere closely related to cryptopine and protopine, and contain the characteristic ten-membered ring-system.

Group III; comprises the strongest bases of the series, these contain free phenohe hydroxyl groups and are easily oxidised by iodine (corytuberme auh group).

Group III comprises alkeloids which belong to the aporphine group (v. Gedamer, Oberhin, and Schoeler, Arch. Pharm. 1925, 263, 61). The numbering of the tetrahydroprotoberberine (Bnck, Perkin and Stevens, J.C.S. 1925, 127, 1462; Gadamer, Spath and Mosettig Ar-h. Pharm. 1927, 265, 675) and the aporphine skeleton is the following :

Group I .- CORYDALINE SUB-GROUP.-This comprises the weak alkaloids of the corydals root, viz. corydaline, dehydrocorydaline, coryhulhine, uscorybulhine, tetrahydropalmatine, and corypalmine. They have in common the property of being capable of oxidation with sodine to the dehydro-compounds which are quaternary hases closely related to berberine and palmatine.

Corydaline was first isolated from C, care hy Wackenroder (Berz. Jahr. 1826, 7, 220), hut it was Freund and Josephi who gave it the correct formula. The yields very con-aderahly, Martindale (Arch. Pharm. 1893, 236, 215) gives 0 9%, Ziegenbein (Arch. Pharm. 1696, 234, 497) quotea 0 57%.

Corydaline (I), G11H17O4N, [a]20 +300 (in ehloroform), m.p. 135°, crystallises from EtOH in colourless, short, six-aided prisms, sparingly soluble in cold, but easily soluble in hot EtOH, Et.O, GHCl, and benzene, (I) contains 2 asymmetric carbon atoms; it is slowly oxidised by atmospheric oxygen to the vellow dehydrocorydaline (11). For absorption spectra of (I) are Dohlie and Lauder (J C.S. 1903, 83, 605), Dohlie and Fox (J C.S. 1914, 405, 1639), and Girardet (J.C.S. 193L 2630). (1) gives two methiodides: one being crystalline, in p. 238-240°, [a]p +1334°, the other amorphous, in p. 64-63°, [a]p +110° (von Bruchhausen and Stippler, Arch. Pharm-1927, 285, 152).

The salts of (1) are crystalline: $B HCI+2H_1O$, columnar eryetals from H₂O, mp. 206-207; B H₁, pale yellow priams of indefinite mp.; B HNO₃, tablets from EtOH, mp. 198, sparingly soluble in H.O; B.HGNS columns from EtOH, mp. 208°; platinichloride, brown crystals, m.p. 227°; the ethyl sulphate forms characteristic, large prisms, m.p. 152 5°.

By exidation of (1) with iodine (Ziegenbein,

Arch. Pharm. 1896, 234, 505), hromine (Martin-dale, Arch. Pharm. 1898, 236, 238) or very dilute

HNO₃ (Dobbie and Marsden, J.C.S. 1897, 71, 657) dehydrocorydaline (II) is obtained, which is identical with a natural base isolated from C. cava, vernyi, ambigua, and solida. (II) is a yellow crystalline powder, m.p. 112-113° (decomp.), the salts are crystalline, B·HCl, 4H₂O, yellow leaflets, soluble in H₂O and EtOH; B·HBr, yellow needles, m.p. 126°; B·HI, 2H,O, small yellow needles; B.HAuCl4, redbrown needles from EtOH-HCl, m.p. 219°; yellow B₂·H₂PtCl₆6H₂O, yellow needles from EtOH—HCl; (II) in solution is a quaternary ammonium base. In the free state it melts and decomposes at 112-113°, has the composition C22H25O5N and must be regarded as a 4-base with the keto-composition containing the groups ·CMeO and NH < instead of ·CMe: N·(OH) < for it forms an oxime, orange-yellow crystals, 165° (Haars, Arch. Pharm. m.p. 243, 171), it also unites with 1 mol. of CHCl₃ to form a colourless crystalline compound m.p. 162-163° and an amorphous acetone compound

(Ziegenbein, ibid. 1896, 234, 492).

By reduction of (II) with zinc and sulphuric acid r-corydaline (III) m.p. 134-135° and r-mesocorydaline (IV) m.p. 158-159°; m.p. 163-164°: Späth and Posega, Ber. 1929, 62, [B], 1029) are formed, which are both optically inactive (Gadamer, Arch. Pharm. 1902, 240, 19; ibid. 1916, 254, 295). (IV) may be resolved into d- and l- forms by crystallisation of the d-camphorsulphonate, the d-form is not identical with natural (I) (Gadamer, ibid. 1902, 260, 50 Haars, l.c.). (III) has not been resolved, but its sulphonic acid can be separated into d- and l- forms by crystallisation of the brucine salts, the d- form is identical with the sulphonic acid prepared from natural corydaline (Gadamer, ibid. 1916, 254, 295).

Insight into the constitution of (I) has principally been obtained by oxidation experiments (cf. Dobbie and co-workers, J.C.S. 1897, 71, 657; 1902, 81, 145; Haars, l.c.). Oxidation with permanganate yielded hemipinic, m-hemipinic acid and corydaldine; HNO3 gave rise to the intermediate oxidation products corydic, corydilic, and methylpyridine carboxylic acids respectively, and the ultimate products were pyridine-2:3:4:5-tetracarboxylic acid, hemipinic and m-hemipinic acids. Important contributions towards the final elucidation of the structure of (I) were made by Spath and Lang (Bcr. 1921, 54, [B], 3074), Gadamer and von Bruchhausen (Arch. Pharm. 1921, 259, 245), Lawson, Perkin and Robinson (J.C.S. 1924, 631), and by von Bruchhausen and Stippler (l.c.). Cf. also Koepfli and Perkin (J.C.S. 1928, 2989) where short survey is given. The following constitution for (I) has been established:

OMe

CH₃

OMe

OMe

OMe

OH

CH₂

OH

II.

Synthetical experiments were carried out by Pictet and Malinovski (Ber. 1913, 46, 2688), Spath and Lang (l.c. 1921), and Spath and Mosettig (Annalen, 1923, 433, 138), but they failed to give a product identical with (III) or (IV). F. von Bruchhausen (Arch. Pharm. 1923, 261, 28) was able to introduce the C-methylgroup into position 13, and he obtained (III) melting at 135-136°; Spath and Holter (Ber. 1927, 60, [B], 1891) obtained synthetic (III) and (IV) in connection with their work on corycavine. An interesting synthesis of (III) and (IV), starting from papaverine, was described by Spath and Kruta (Ber. 1929, 62, [B], 1024).

Corybulbine and isoCorybulbine.—Corybulbine was first isolated by Freund and Josephi (Annalen, 1893, 277, 1); isocorybulbine was found by Gadamer and Ziegenbein (Arch. Pharm. 1902, 240, 634). They occur in corydalis roots in an amount of 0.05% (Gadamer, ibid. 24), and 0.185% respectively (Bruns, Arch. Pharm. 1903, 241, 634). The yields vary considerably. Makoshi (Arch. Pharm. 1908, 246, 381) isolated (I) from a Korean corydalis, possibly C. ambigua (see also Chou, A. 1929, 1085). Both alkaloids (I) and (II) are sensitive to light.

Corybulbine (V), C₂₁H₂₅O₄N, [a]_D +303·3°, m.p. 239-240°, crystallises from boiling EtOH in colourless needles, sparingly soluble in EtOH, Et₂O, and CH₃·CO₂Et, soluble in CHCl₃, acetone, or hot benzene. (V) is a weak base.

B·HCl, colourless prisms, m.p. $245-250^\circ$, slightly soluble in H_2O ; platinichloride and aurichloride are amorphous. By oxidation with iodine optically inactive dehydrocorybulbine, $C_{21}H_{21}O_4N$ (dark violet needles, from H_2O , m.p. $175-178^\circ$), is formed.

By reduction an optically inactive r-corybulbine is obtained, m.p. 220-222°, which has not been resolved with bromocamphorsulphonic acid (Bruns, Arch. Pharm. 1903, 241, 650).

isoCorybulbine (VI), C₂₁H₂₅O₄N, [a]_D+299·8°, m.p. 179–180°, crystallises from EtOH in colourless leaflets more soluble in EtOH than (V); by this property (VI) may be separated from (V). (IV) is a tertiary base with 1 OH and 3 OMe-groups. For colour reactions of (I) and (II), see Gadamer (ibid. 1902, 240, 52). (II) is oxidised with iodine to dehydroisocorybulbine, which yields rac. isocorybulbine, m.p. 165–167° on reduction. Dobbie, Lauder and Paliatscas methylated the phenolic OH-group in (V) and obtained corydaline (J.C.S. 1901,79, 87), also (VI) on methylation yields corydaline (Spath and Dobrowski, Ber. 1925, 58, [B], 1274). All 3 alkaloids give the same apocorydaline (hydrochloride, m.p. 220–222°) on demethy-

lation (Bruns, 1c.) This shows that (V) and 1911, 249, 30). (I) and (II) have both the same (VI) are closely related to corydalme which is a methyl ether of (V) or (VI). Spath and Dobrowski (Ic) succeeded in demonstrating the exact positions of the phenolic groups and both alkaloids were finally obtained by partial demethylation of corydaline (Spath and Holter, Ber. 1926, 59, [B], 2800), (V) having the phenolic OH-group in position 3 and (VI) in position 2 (cf. structure of corydaline).

d.Tetrahydropalmatine Was isolated Spath, Mosettig, and Trothandl (Ber. 1923, 56, 877) from C. cava grown near Vienna. yield from dried corydalis roots is about 0 4% the extraction was effected by the method of Gadamer (Arch. Pharm. 1902, 240, 21). Separation of phenolic from non-phenolic hases was achieved hy nsing caustic soda, d-tetrahydropalmatine was separated from corydaline by the insolubility

of its hydrochloride.

Tetrahydropalmatine (VII), C21H25O4N, [a] +292 5° (in EtOH), m p. 142°, erystallises from dilute EtOH, is easily soluble in organic solvents, except Et,O and light petrolcum, On exposure to air it is coloured yellow; by on daton with rodine, the quaternary rodide of palmatine (mp. 238-239°) is obtained. Spath and Mosettig (Ber. 1926, 59, [B], 1496) on the ased d tetrahydropalmatine atarting from d-tetrahydroberbenne (vv.), removing the methylenedioxy group with phloroglueinol sul-phurie acid and methylating the phenoho base. The product obtained was identical in every respect with natural (VII).

For constitution of (VII), of palmatina (Cal-

UMBA ALEALOIDS).

Corypalmine (VIII), CacHasO4N, mp 235-236°, [a]14 +280° (m CHCl3), was first isolated hy Spath, Mosettig, and Trothandl (Ber. 1923, 56, [B], 875) in a yield of about 0 005%. It forms small white crystals and is sparingly soluble in MeOH. By methylation of the I phenolic OHgroup with diazomethane, d-tetrahydropalmatine q.v.) ia obtained. In a later paper (Ber, 1925, 58, [B], 2133) Spath and Mosettig demonstrated the exact position of the OH group in (VIII) and proved it to be d-tetrahydro jatrorrhizine (q v. calumba alkaloids). (VIII) was synthesised Spath and Mosettig (Ber. 1927, 60, (B1, 383) from d. tetrahydroberberine, the metbylenediox, group was replaced by OH groups with phloroglucinol H2SO4 and the resulting pheno-lic base was partially methylated; from the mixture of products thus obtained, (VIII) could be isolated. The phenolic OH group takes the same position 3 as in coryhulbine (q v.).

There is evidence for the existence of secony palmine in C. cara, differing from corypalmine by the attachment of the phenolic OH-group to C.atom 2 (d.tetrshydrocolumbamin), cf. Gadamer, Spath and Mosettig (Arch. Pharm.

1927, 265, 675).

Group 11 .- CORYCAVINE SUB GROUP .- This group comprises the three alkaloids corycavine (I), corycavamine (II), and corycavidine (III).
(III): R_k+R₀=CH₂; R₃=R₄=OMe

(Annalen, 1893, 277, 1), (11) by Gadamer, Ziegenbein, and Wagner (Arch. Pharm. 1902 The three elkaloids do not react with kelo-240, 19), and (111) by Gadamer (Arch, Pharm, reacenta,

structure, (I) is the racemic form, (II) is optically active and is easily transformed into (I) by beating to above the melting point (see F. von Bruchhausen, Arch. Pharm. 1925, 263, 601).

Corycavine (1), C21H31O5N; cf. Gadamer and von Bruehhausen (Arch. Pharm. 1922, 280. 97) where earlier references are given ; it occurs in C. tata in amounts from 0 07% (Legerlott, Arch. Pharm. 1918, 258, 161) to 0 11% (Gadamer and von Bruchhausen, sbid. 161). (1) crystal lises from EtOH or Et₂O in rhombic tablets. m p. 216-218°, insoluble in cold EtOH and H.O. BHC! needles (from H2O), mp. 219'; BHBr quadrangular flat plates from HO;

B methiodide rhombic tablets, m.p. 218°, Resolution of (I) into d. and I. (II) has not yet

been effected,

Corycavamine (II) occurs in an amount of 0 3% (Gadamer, Arch. Pharm, 1902, 240, 84) It erystallises in rhombic columns from ether. alcohol, or CH, CO, Et, m p. 148-149", [a], +166° (in CHCl3), transformation (II) -> (I), see above. The salts except the sulphate crystallise very easily, the nitrate is speringly soluble and is used to purify (II); B HCl, small needles, soluble in H₂O; B HBr speringly soluble needles; B HI, yellow voluminous,

aparingly soluble needles.

Corycavidine (III), C₃₃H₂₅O₅N, occur is an amount of 0 05%, [a]_b +203 I⁵ (in CHCl₃), m p. 212-213, crystalises from CHCl₂-E10H in transient, colourless crystals with I mol. of CHCl, practically insoluble in EtOH sid EtO. Crystalline hydrochloride and springly abluble intrate. By continued heating at 20' (III) is transformed into an optically inactive form, melting at 193-195°

Pseudocorycavine (see Gaebel, Pharm. 1910, 246, 249) 19 a mixture of (I) and (III) (cf. Gadamer, Arch. Pharm, 1911, 249, 231) which is difficult to separate, but it can be effected by recrystallisation of the hydrochlorades.

Subject to a slight modification of (I) by Spath and Holter (Ber, 1927, 60, [B], 1891), the constitutions of (I), (II), and (III) have been conclusively proved by F. von Bruchhausen (Arch. Pharm, 1925, 263, 570),

(III):
$$R_1 + R_9 = CH_{\frac{1}{2}O}$$
; $R_3 = R_4 = OMe$

little importance (cf. Gadamer, Arch. Pharm. | MeO groups in (I) and (III).

1905, 243, 147).

Group III .-- CORYTUBERINE SUB-GROUP. Besides bulbocapnine (q.v.) this group comprises the closely related phenolic aporphine alkaloids corytuberine (I), corydine (II), and isocorydine (III). (I) and (II) occur in C. cava (0.7-1.5% and 0.3% respectively). (III) and the lavo-form of (II) were found in a Korean corydalis by Go (Amer. Chem. Abstr. 1930, 24, 620); (II) and (III) have been isolated from Dicentra canadensis and D. oregana (Manske, Canad. J. Res. 1932, 7, 258; 1934, 10, 765). Dicentra formosa contains 0.02% of (I) and 0.03% of (II) (Manske, Canad. J. Res. 1934, 10, 521).

For preparation of (I), cf. Gadamer, Arch. Pharm. 1911, 249, 645; for preparation of (II), Arch. Pharm. 1902, 240, 96.

Corytuberine (I), $C_{19}H_{21}O_4N,5H_2O$, $[a]_D^{20}$ +282.6° (in EtOH), m.p. 240°, crystallises from hot H₂O in silky needles, soluble in EtOH slightly soluble in CHCI3 and CH3 CO2Et, insoluble in benzene and Et2O. The salts are crystalline but somewhat difficult to prepare, because (I) is a weak base and probably forms a phenolbetaine (Gadamer); the hydrochloride is sparingly soluble in H₂O.

(II) and (III) are obtained by methylation of (I) with diazomethane; Gadamer experienced great difficulties in completely methylating (I); he only succeeded by doing it with diazomethane in statu nascendi in amyl ether solution (l.c. 1911).

Corydine (II), $C_{20}H_{23}O_4N$, $[a]_D + 204\cdot4^\circ$ (in CHCl₃), m.p. 124-125° from dilute EtOH with $\frac{1}{2}$ mol. of EtOH or m.p. 149-150° (dry). It is readily soluble in CHCl₃ and EtOH, soluble in Et₂O. Methiodide, m.p. 190-191° (228-230° according to Manske, l.c. 1932), $[a]_D^{20} + 154\cdot6^\circ$, easily soluble in H_2O . By indicate the part of the contraction of iodine (II) is oxidised to the quaternary dehydrocorydine iodide (m.p. 248°); subsequent reduction yields racemic corydine, m.p. 168°, which can be resolved with tartaric acid into d- and l-forms (l-corydine, m.p. 151° , $[a]_{D}^{26} - 206^{\circ}$).

(II) HCl, m.p. 258°, is sparingly soluble in H₂O; separation from the hydrochloride of (III)

is thus possible.

isoCorydine (III), C₂₀H₂₃O₄N, [a]_p +195·3° (in CHCl₃), m.p. 185°, crystallises in 4-sided tablets from CH₃·CO₂Et-MeOH. (III) is less soluble in Et₂O than (II); (III)·HCl is more soluble in H₂O than (II) ·HCl. Methicdide, m.p. 213-214° (decomp.), [a]_p²⁰ +143·3°, speringly soluble in H₂O. No experingly soluble in H₂O. sparingly soluble in H2O. No crystalline product was obtained by oxidation with iodine solution. For differences in colour reactions cf. Gadamer (l.c. 1911, 676)

Gadamer (l.c. 1911, 641, 669) on the basis of his investigations assigned structures to (I), (II), and (III) which had to be modified later in minor details. By synthesis of (I) dimethylether (Gulland and Haworth, J.C.S. 1928, 1834, and Spath and Hromotka, Ber. 1928, 61, [B], 1692) it was demonstrated that (I) belonged to the aporphine alkaloids. Späth and Berger (Ber. 1931, 64, [B], 2038) synthesised corydine starting from bulbocapnine (q.v.) and conclusively

Pharmacologically the three alkaloids are of | demonstrated the arrangement of the OH and

(I): R_1 and R_4 =MeO; R_2 and R_3 =OH. (II): R_1 , R_2 and R_4 =MeO; R_3 =OH. (III): R_1 , R_3 and R_4 =MeO; R_2 =OH.

RECENTLY ISOLATED CORYDALIS ALKALOIDS -(a) Non-phenolic bases.—From C. cava grown near Vienna, Späth and Julian (Ber. 1931, 64, [B], 1131) isolated d-tetrahydrocoptisine, [a]D +310°, m.p. 203-204° (for constitution, see COPTISINE), d-canadine, [a]_D¹⁵ +299°, m.p. 131-132° (for constitution, see Berberine) and hydrohydras-

tinine, m.p. 66°.
(b) Phenolic Bases.—Gadamer, Späth and Mosettig (Arch. Pharm. 1927, 265, 675) isolated the 2 alkaloids: d-tetrahydrocolumbamine, m.p. 240-241° (for constitution, see COLUMBA-MINE) and 2-9-dihydroxy-3-10-dimethoxytetrahydroprotoberberine (for numbering, cf. General Introduction to this article), m.p. 195°, which on complete methylation with diazomethane yielded d-tetrahydropalmatine. new bases isolated from Corydalis ambigua, see Hnang-Milnon (Ber. 1936, 69, [B], 1757).

Schl.

CORYDINE v. Corydalis Alkaloids, CORYTUBERINE SUB-GROUP.

CORYNANTHINE, one of the Yohimboa alkaloids.

CORYPALMINE v. CORYDALIS ALKALOIDS, CORYDALINE SUB-GROUP.

CORYTUBERINE v. CORYDALIS ALKAloids, Corytuberine Sub-Group.

COSTENES. The sesquiterpene hydrocarbons, a and β-costenes, C₁₈H₂₄, were isolated by Semmler and Feldstein (Ber. 1914, 47, 2692) from the roots of Saussurea Lappa. These hydrocarbons, the homogeniety of which is doubtful, have the following constants: a-, b.p. 122-126°/12 mm., d^{21} 0.9014, $n_{\rm D}^{21}$ 1.49807, [a]_D -12°; β -, b.p. 144-149°/18 mm., d_{22} 0.8728, $n_{\rm D}^{22}$ 1.4905, [a]_D +6°. No crystalline derivatives have been prepared.

COSTOL. The primary sesquiterpene alcohol, costol, $C_{15}H_{24}O$, b.p. $169-171^{\circ}/11$ mm., d^{21} 0.9803, $n_{\rm D}$ 1.5200, $[a]_{\rm D}$ +13°, was separated by Semmler and Feldstein (Ber. 1914, 47, 2687) from the oil from the roots of Saussurea lappa. The alcohol, which is probably dicyclic, gives on oxidation with chromic acid an aldehyde, $C_{15}H_{22}O$, b.p. $164-165^{\circ}/15$ mm., d^{22} 0.9541, $n_{\rm D}$ 1.50645, $[\alpha]_{\rm D}$ +24°, semicarbazone, m.p. 217-218°.

J. L. S.

COSTUS, ESSENTIAL OIL OF. The phenone with less than one molecular proportion oil distilled from the root of Squasurea lappa of diazomethane et -12° . The synthesis, how Clarke (Aplotants lappa Decaisne) indigenous to ever, does not establish the orientation of the the mountain slopes of the Kashmir valley. methoxyl group in cotoin. The root is used for incense and to preserve the celebrated Kashmir shawls from the attack of moths. The oil is used in perfumery Yield 1-3%.

Constituents .- Costic acid, costolactone, hydrocostolactone, the sesquiterpene alcohol, costol (q v.), a hydrocarbon aplotaxene and the two sesquiterpenes-a-costene and & costene (qv), with traces of phellandrene end camphene.

Characters.- A light yellow oil, spgr 0 982-0 987, optical rotation about +15°, n21 1 510-1.520.

CTB COTARNINE, an oxidation product of narcotine, an opium alkaloid.

COTO BARK. True coto bark is obtained from Bolivia, the Amazons (Brazil), and Venezuela, It has been considered identical with Winter's bark [from Dramys Winters Forst,], but its source is really unknown (Hesse, Annalen, 1894, 282, 191) It comes into the market in long strips about 8 cm, broad end 12 mm, thick, and is tough and hard. The colour is cinnamon brown, and a cross section shows numerous yellow spots It has an anoms numerous years aromatic odour, and contains cotour, dicotour, hydrocotoun, and methylhydrocotoun, resuns, tan nin, etc. The supply of the true coto bark is limited, and inferior bark's, chiefly para coto bark, are sold; these contain protocotoin and methyl protocotorn, but little or no cotom.

Cotorn can be molated from true coto bark by extracting the powdered bark with ether, removing the ether, and mixing the residue, whilst still hot, with light petroleum. A resinous oily mass separates from which the solution of cotoin is decanted Cotoin forms yellow prisms, mp. 130-131°, is difficultly soluble in cold water, readily soluble in hot, is fairly soluble in alcohol, ether and chloroform, but sparingly soluble in light petroleum or benzene. It dissolves in alkalis forming yellow solutions from which it is precipitated on acidificotion, it yields phloroglusinal and benzoid acid as decomposition products.

Cotom has been the subject of much investigation, by Ciamician and Silber (Ber. 1891, 24, 299, 2977; 1892, 25, 1119; 1893, 26, 777; 1894, 27, 409, 841; 1895, 28, 1549), Jobst N. Repert. Pharm. 1876, xxv, 23; Ber., 1878, 11, 1031), Johnt and Hesse (Ber. 1877, 10, 249 : Annalen, 1879, 199, 17), and Hesse (I.c., Ber. 1893, 26, 2790; 1894, 27, 1182). As a result it has been shown to be 2.6 dihydroxy-4. methoxy benzophenone.

Spith and Fuchs (Monatah., 1921, 42, 267)
obtained cotoin, hydrocotoin (cotom methyl ether), and a small amount of cotom dimethyl ether hy methylsting 2.4.6 trihydrocybeneomatter, mp. 144". Methylation of the former

From a consideration of the products obtained by methylation and ethylation of cotom, Spath and Wessely (Monatsh. 1928, 49, 229) conclude that the methoxyl group of cotoin is in the para position. The properties of the derivatives of cotom obtained by p toluenesulphonylstion (Karrer and Lichtenstein, Helv. Chim. Acta, 1928,

11, 789) also point to the same conclusion.

Cotom has been used pharmaceutically in cases of diarrhora and of phthisis.

The following derivatives of cotoin have been prepared · Discetylcotion, prisms, m.p. 94°, is pro-

duced by the action of acetic anhydride. If sodium acetate is used in the acetylation, the diacetyl derivative is accompanied by methoryhydroxyphenylcoumarın acetate, m.p. 142°;

(Jobst and Hesse, Annalen, 1879, 199, 28; Ciamician and Silber, Ber. 1894, 27, 419) Benzoylcotosn, prisms, mp. 110-112°, and dibenzoylcotosn, needles, m.p. 134-135° (Hesse, Annalen, 1894, 282, 194)

Dimethylcotoin, mp. 138°, obtained by the action of methyl locade and alkali on cotoin, yields a monoacetyl derivative, colourless needles, m p 150°, and hence probably has the formula :

HO-C.H-Me(OMe). CO C.H.

(Ciamician and Silber, Ber. 1894, 27, 418). Dibromocotion, colourless needles, m p. 1161, coton oxime, C14H15NO4, plates, soluble in alcohol, difficultly soluble in water (Ciamician end Salber, Ber. 1894, 27, 415)

Mononstrosocotom, dark red lesflets or orange-yellow needles, m p. 153-154° (Pollak, Monatsh. 1901, 22, 996).

Cotoinazzo-o-toluene, orange-yellow needles, m.p. 203-204°, cofoinazo-p toluene, orange-yellow needles, m.p. 207-208°, and cofoinazo-p toluene, orange-yellow needles, m.p. 207-208°, and cofoinazo-p toluene, needles, needle benzene, orange-yellow needles, mp 183-184, diacetylcotomazobenzene, scarlet needles, mp 155-166° (Perkin and Martin, J.C.S. 1897, 71,

Cotom ethyl ether (2 hydroxy 4 methoxy 6 ethoxybenzophenone), m.p. 91-92°, cotom methyl ethyl ether (2.4-dimethoxy 6 ethoxy benzophenone), m p. 103-104°, cotoin diethyl (4-methoxy 2:6 diethoxyhenzophenone). ether m.p 82-83°, bp. 160-170°/0 001 mm. (Spath and Wessely, I c.)

112°, of hydrocotoin (2-hydroxy-4:6-dimethoxybenzophenone) (Karrer and Lichtenstein, I.c.).

A number of polyhydroxybenzophenone derivatives, including products found in coto bark and related to cotoin, have been prepared by W. H. Perkin and Robinson (Proc. Chem. Soc. 1906, 22, 305).

A. G. P. and E. J. C.

COTOIN v. Coto Bark.

COTTON FLOWERS. The flowers of the cotton plant, according to Watt (" Dictionary of the Economic Products of India"), are used as

a dyestuff in the Manipur district.

The colouring matters present in the flowers of the ordinary Indian cotton plant, Gossypium herbaceum, have been studied by Perkin (J.C.S. 1899, 75, 826; 1913, 103, 650), who isolated a mixture of quercetin and gossypetin by digesting a concentrated alcoholic extract of the flowers with boiling dilute hydrochloric acid. The mixture was acetylated and the acetyl derivatives of the colouring matters separated by fractional crystallisation from acetic anhydride, in which acetylgossypetin is less soluble.

Quercetin, the aglucone of the glucoside quercitrin, the colouring matter of quercitron bark, is 3:5:7:3':4'-pentahydroxyflavonol:

Gossypetin, $C_{15}H_{10}O_8$, yellow needles, m.p 311-313°, closely resembles quercetin in appearance, and is readily soluble in alcohol, but only very sparingly soluble in water. Concentrated alkaline solutions dissolve it forming orange-red liquids which, on agitation and dilution with water, become green, then blue, and finally assume a dull brown tint. Alcoholic lead acetate gives a deep red precipitate in the cold, passing into dull brown at the boiling-point, and with alcoholic ferric chloride a dull olive-green liquid

Gossypetin forms crystalline compounds with mineral acids and yields a monopotassium derivativo, $C_{15}H_9O_8K$, with alcoholic potassium Hexa-acetylgossypetin, colourless needles, melts at 228-230°. By fusion with alkali, gossypetin gives protocatechuic acid and a phenolic substance which has not yet been identified.

When gossypetin hexamethyl ether, colourless needles, m.p. 170-172°, is hydrolysed with alcoholic potash, protocatechuic acid dimethyl ether and gossypitol tetramethyl ether, C₁₂H₁₆O₆, needles, m.p. 115-116°, are produced. In a similar manner, gossypetin hexaethyl ether, m.p. 144-146°, yields protocatechnic acid diethyl ether and gossypitol tetraethyl ether,

HO·C₆H(OEt)₃·CO·CH₂·OEt,

m.p. 110-111°. The latter forms an oxime, C₁₆H₂₅O₆N, m.p. 127-129°, and when oxidised with permanganate gives gossypetonic acid,

gives the p-toluenesulphonyl derivative, m.p. | HOC6H(OEt)3 COCOOH, yellow needles, m.p. 154-155°.

As a result, Perkin concluded that gossypetin is hydroxyquercetin:

By the oxidation of quercetin with chromic acid, Nierenstein and Wheldale (Ber. 1911, 44, 3487) obtained quercetone, which, on simultaneous acetylation and reduction and subsequent saponification, yielded hydroxyquercetin. To this substance, which is not identical with gossypetin, formula (I) was assigned, and its synthesis was described by Nierenstein (J.C.S. 1917, 111, 4). On the other hand, Baker, Nodzu and Robinson (ibid. 1929, 74) obtained results which confirm the work of Perkin but are irreconcilable with that of Nierenstein. These authors synthesised gossypetin by veratroylation of the trimethoxyresacetophenone (II). The hydroxypentamethoxyflavone (III) thus formed gave gossypetin (I) on demethy. lation.

II.

Gossypitone.—When an alkaline solution of gossypetin is exposed to air, the blue liquid on acidification yields a reddish-brown precipitate which, on gentle warming, becomes crystalline. Again, if benzoquinone is added to a solution of gossypetin in cold absolute alcohol, a similar reaction occurs. On keeping for a few minutes crystals begin to separate, and by then heating to 50° a semi-solid mass is obtained. This substance, gossypitone, C15H8OH, consists of microscopic needles of a dull red colour, which are difficultly soluble in the usual solvents. It dissolves in dilute alkalis with a pure blue coloration and its solution in concentrated sulphuric acid is dull brown. Sodium bisulphite solution reconverts it into gossypetin. Gossypitone possesses strong dyeing properties, and gives the following shades on mordanted woollen

Aluminium . . Orange-brown. Chromium . Dull brown. Tin Orange-red. Iron . . Deep olive.

These shades are identical with those given in | and gives with lead acetate a deep red precipities circumstances by geosypetin itself, and it tate. It is almost insoluble in boung acctore, is accordingly evident that during the dyeing a property which readily distinguishes it from operation oxidation of the latter to gossypitone

takes place.

The glucosides present in cotton flowers have been examined by Perkin (J.C.S. 1909, 95, 2191), whn in the first place employed the Egyptian variety. The Egyptian cottons appear, according to the researches of Messrs, Fletcher and Balls, to be derived from natural crosses of brown Peruvian cottons with the Sea Island variety.

Three glucosides, queremeritrin, gossyputrin, and isoquercitrin were isolated, with difficulty, from a concentrated alcoholm extract of the

Quercimeritrin, C21H20O12,3H2O, small, ghstening, hright yellow plates, m p. 247-249°, is insoluble in cold and fairly readily soluble in boiling water. Its alkaline splutione possess a deep yellow tint ; with aqueous lead sectate it gives a bright red precipitate, and with ferric chloride an olive-green coloration. Octa-acetylquercimentrum forms needles, m p. 214-216°. By hydrolysis with dilute sulphurio acid quercimentrin yields quercetin and glucose

On mordanted woollen cloth, quercimentria

gives the following shades :

. Orange yellow. Aluminium . Chromsum . Reddish brown. Tin . . Bright orange Iron . Ohve-brown.

With the exception of the iron mordanted pattern, which is of a rather browner character. the colours thus produced closely resemble those given by querectin itself when dyed in a similar manner, and are widely different from those

given by rutin and quereitms.

The position of the sugar group in queremeritrin has been determined by Attree and Perkin (J.C.S. 1927, 234). With diazomethane the glucoside yields almost colourless needles of the pentamethyl ether, CalH15O, (OMe), 2H1O, m.p. 203-205°, sintering at 197°. By hydrolysis, colourless needles of a sparingly soluble quercein tetramethyl ether, m.p. 284-285°, ore obtained, the acetyl derivative of which melts at 174-176°. As this, when digested with alcoholic potash at 180°, readly gives versing acid and phloro-glucinol monomethyl ether, it is evidently the hitherto unknown 3.5.3'.4'.tetramethoxy.compound. Quereimentrin, therefore, has the formula : OH

which is in harmony with its dyeing properties, Onereimeritrin is also present in the flowers

of the Prunus emarginata ? (Finnemore, Pharm. J. 1910, (iv), 31, 604) and Helianthus annuus (Sando, J. Biol. Chem. 1926, 68, 407).

Gossypitrin, C₂₁H₂₀O₁₂, yellow needles, m p. 240-242°, (+H₂O) m.p. 200-202°, dissolves in alkalis with a pale yellow coloration, usually now treated as one species but originally

a property which readily distinguishes it from querementrin, which is fairly soluble in this solvent. When bydrolysed with dilute sol. phuris acid, gossypitrin yields gossypetin and glueose. Acetylgossypitrin,

colourless needles, m.p. 226-228°, is almost meduble in alcohol.

Gossypetrin reacts, like gossypetin itself, with benzoquinone and forms a quinone, gossypitrone. C₂₁H₁₀O₂₃, maroon-coloured needles, which possess an definite melting-point although fusion of the product occurs about 255-259°. By the action of warm dilute sulphurous acid solution it is reconverted into gossypitan.

Identical shades are given by gossypitrin and gossypitrone on mordanted wnol:

Aluminium . . Dull yellow. . Reddisb brown. Chromsum . Tin . . Bright orange, . Dark olive brown.

and it is thus evident that the former is con verted into the latter during the dyeing process

vertea mot tee atter during me uyeng process. The engar residue of gosyntim is probably attached to the hydroxyl group in the 7-pouton is Querclitin, Cyll 10, 22 H, O, pale yellow needles, m. p. 217-218", is sparingly soluble in water and dissolves in alkaline solutions with a deep yellow tunt. It gires a highly silow precipitate with aqueous lead a coctate and when hydrolysed yields quercetin and glucose.

By methylating isoquercitrin with diszo-methane, Attree and Perkin (i.e.) obtained pale yellow, flat needles of a methylated derivative, m.p. 150-152°, which on hydrolysis gave 5.7:3':4'-tetramethoxyflavonol. isoQuercitrin

thus has the structure :

OH

On mordanted wool isoquercitrin gives the following shades which are entirely distinct from those given by quereimeritrin and, although alightly paler, resemble those yielded by quercitrin :

Aluminium . . . Golden yellow. . Brownish yellow. Chromium . . . Lemon yellow. . Brownsh obve. Iron .

isoQuercitrin has also been isolated from hrown-husked maize (Sando and Bartlett, J Binl. Chem. 1922, 54, 629) and from mulberry leaves (Oku, J. Agric, Chem. Soc. Japan, 1934. 10, 1029).

Among the types of cotton flowers there are red, pink, yellow and white flowered plants Perkin (J.C.S. 1916, 199, 145) examined the following types: red flowered, G. arborum Line. ; pink, G. sanguineum Harsk ; yellow and white, two varieties of G. neglectum Tod,

described as G. neglectum and G. rosarum. As a result it was found that the red flowers of G. arboreum contain isoquercitrin, whereas in the yellow flowers of G. neglectum, gossypitrin and isoquercitrin were present. On the other hand, the white flowers of G. neglectum and the pink flowers of G. sanguineum gave amounts of colouring matter too small for complete identification, although the respective products obtained resembled in their properties apigenin and

An examination of the ordinary Indian cotton flower, G. herbaceum, available only in small amount, showed the presence of gossypitrin and isoquercitrin. On the other hand, Neelakantam, Rao and Seshadri (Proc. Indian Acad. Sci. 1935, 2, A, 490) find that the petals of G. herbaceum from Coimbatore, South India, contain gossypitrin, quercitrin (instead of isoquercitrin), and a flavonol glucoside herbacitrin, m.p. 247-249°, with small quantities of gossypetin. Herbacitrin forms an octa-acetyl derivative, m.p. 214-216°, and on hydrolysis yields herbacetin, m.p. with 3:5:7:8:4'-penta-279-281°, identical hydroxyflavone synthesised by Goldsworthy and Robinson (J.C.S. 1938, 56). The colour reactions of herbacitrin indicate that it is the 7-glucoside of herbacetin (Neelakantam and Seshadri, Proc. Indian Acad. Sci. 1937, 5, A,

According to Neelakantam, Rao and Seshadri (ibid. 1935, 1, 887), quercetin and quercimeritrin are present in the flowers of G. hirsutum in proportions which vary with variety, locality and season. From the petals of *G. indicum*, Ncelakantam and Seshadri (*ibid.* 1936, 4, A, 54) isolated gossypin, m.p. 230° (decomp.), which by hydrolysis yielded gossypetin and glucose and a non-glycosidio substance, C₁₆H₁₂O₇, m.p. 255-257°, the acetyl derivative of which melts at 186-187°. In addition, gossypitrin, herbacitrin, and quercetin were isolated.

Gossypetin is also present in the flowers of the Hibiscus sabdariffa or "red sorrel" of the West Indies, a small shrub which is widely cultivated throughout the hotter parts of India and Ceylon (Perkin, J.C.S. 1909, 95, 1855). The stems yield the "Rozelle hemp" of commerce.

A. G. P. and E. J. C. COTTONSEED, COLOURING MAT-Cottonseeds contain, in addition to cottonseed oil, a highly toxic phenolic substance, gossypol, which romains dissolved in the oil when this is expressed.

By extracting the oil with caustic soda and neutralising the extract, a voluminous precipitate of the colouring matter admixed with fatty acids and the products of its own oxidation separates. Marchlewski (J. pr. Chem. 1899, [ii], 60, 84) isolated gossypol from a partially purified preparation of this character by extraction with ether and repeated crystallisation from acetie acid. The small yellow prisms melted at 188°, and dissolved in sulphuric acid with a cherryrcd coloration, a reaction by which the presence of gossypol in cottonseed oil can be detected. Gossypol dissolves in alkalis forming yellow solutions which are readily susceptible to air slowly fade. An alcoholic solution of gossypol

gives with ferric chloride a dark green coloration and with neutral and basic lead acetate a deep yellow precipitate. According to Marchlewski, gossypol appears to possess 2 hydroxyl groups, but contains no methoxyl groups, and is not a glucoside. It dyes iron-mordanted material a grey shade, and can be employed on cotton as a mordant for the basic colouring matters.

Carruth (J. Amer. Chem. Soc. 1918, 40, 647) isolates gossypol from the decorticated cottonseed kernels which have been crushed by rollers before they are "cooked" in the oil mill. From these a considerable amount of oil is removed by percolation with light petroleum, and the residue is extracted with ether and tho extract concentrated. The product is treated with acetic acid and, on keeping, crystals of gossypol acetate separate. To obtain gossypol itself, an ethereal solution of the acetate is treated with water and the ether distilled off. The gossypol thus remains suspended in the

water as crystalline crusts.

A second interesting method consists in adding aniline to an ethereal extract of cottonseed. The solution, after warming on the water-bath and standing, gradually deposits an orangeyellow microcrystalline precipitate which apparently consists of an aniline salt of gossypol. The yield is practically quantitative, and the process may be used to estimate gossypol in extracts of cottonseed products. To regenerate gossypol, the aniline compound is dissolved in hot alcoholic potassium hydroxide, the aniline removed by steam distillation, and the gossypol precipitated by acidifying the alkaline liquid.

Gossypol has been examined by Clark (J. Biol. Chem. 1927, 75, 725; 1928, 76, 229; 77, 81 78, 159; J. Amer. Chem. Soc. 1929, 51, 1475, 1479), who isolated the substance by the method of Carruth (l.c.). After removal of acetic acid the product separated from a mixture of ether and light petroleum in crystalline form, m.p. 214° (decomp.). Analysis gave the formula $C_{30}\hat{H}_{30}O_8$. Gossypol yields a dioxime, m.p. 315°, and by acetylation a hexa-acetyl derivative, m.p. 276-277°, four of the acetyl groups of which are readily hydrolysed, whilst the remaining two are more resistant to attack. Gossypol thus contains 2 carbonyl groups and 6 hydroxyl groups.

Gossypol acetate, $C_{30}H_{30}O_8$ · $C_2H_4O_2$, m.p. 189–190° (decomp.), a molecular compound of gossypol and acetic acid, is evidently the substance described by Marchlewski as the free colouring matter. It is a comparatively stable substance, and when heated does not evolve acetic acid readily below 180°. The absorption curve shows maxima at 3,780 Å and 2,390 Å and a minimum at 3,220 Å (Grünbaumówna and Marchlewski, Biochem. Z. 1936, 286, 295).

When heated at 215°, gossypol loses 2 mols. of water and forms anhydrogossypol, m.p. 268°

after softening at 229-230°.

With aniline, gossypol forms a dianilide, m.p. 302-303°, apparently a condensation product of 1 mol. of gossypol with 2 mols. of aniline and elimination of water, but not a salt of gossypol as Carruth (l.c.) has suggested.

According to Karrer and Tobler (Helv. Chim. oxidation and rapidly become violet and then Acta, 1932, 15, 1204), gossypol, m.p. 199° yields a quinoxaline derivative, m.p. 298° (after

sintering and decomposition), and is thus an mattern of 4 quinone groups without loss of a diketone or, more prebably, an o quinone The dihydrazone derivative, mp. about 285°, gives with sodium ethoxide at 180° a aubstance, CasHa4O4Na. which contains the grouping

Two of the hydroxyl groups of gossypol titrate as acids, and two are resistant to hydrolysis when esterified.

Oxidation of gossypol with cold alkaline permanganate yields formic, acctic and isobutyric acids. The yield of the last named is approximately one molecular proportion, and this is considered to indicate the presence of an isobutyl group in the eide-chain of gossypol With chromic acid, hexa-acetylgossypol gives a bright yellow substance, tetra acetylgossypolone,

which chars at 230°. This reaction suggests the removal of CaHa from, and the formation of two observes that during the manufecture of cotton additional quinone groups in, the gossypol nucleus (Clark, lc). With ozone, gossypol yields gossypolic acid, probably (C₁₂H₁₄O₄)_p m p 241°, which is shown by colour reactions

Schmid and Margulies (Monatsh. 1935, 65, 391), by catalytic reduction of gossypol, ohtained hydrogossypol, C30H34Og, decomposing at 320' which does not form an additive compound with acetic acid but yields a here ecetyl derivative (decomp 235°) on acetyletion. When distilled with zine dust it gives & isoamylnaphthalene and s substance contenning oxygen, m p 234-238°, hp. 180°/12 mm By oxidation of hydro gossypol with alkaline peroxide, butyrie acid is obtained together with unidentified com

By extracting cottonseeds with chloroform, Podolskoya (Biochem Z. 1936, 284, 401) isolated a red crystalline gossypol which differs from yellow gossypol in its absorption epectrum. On the other hand, both yield the same aniline derivative, and the red pigment changes to the yellow in alcoholic solution. It is suggested that the red colouring matter is present in the seed, and is changed to the yellow under the influence of reegents.

apoGossypol.-By heating gossypol with 40% sodium hydroxide at 100°, apogossypol, C₁₈H₃₀O₆, is obtained together with formic acid resulting from the elimination of the 2 yields a hexamethyl ether, m p. 259°, and a hexa

acetyl derivative, m.p. 291°.

Oxidation of hexa ecetylapogossypol in acetic acid with chromic acid yields tetra acetylapogossypolone, C, H, O, (OAc), m.p. 230°. apocostypole or another the southerd in a southerd in a sum are manner to yellow the cost of the sum are manner to yellow the cost of the sum are manner to yellow the cost of the sum are manner to yellow the cost of the sum are manner to yellow the cost of the sum are manner to yellow the cost of the sum are manner to yellow the sum are made and the sum are the sum a

carbon from the apogossypol ekeleton,

B- and C-Gossypols .- When gossypol is heated to 186-190° it fuses with the evolution of vapour, and then solidifies to a black mass On treating the residue with ether, a sparingly soluble yellow crystaline compound, B-gossypol separates, which is much less toxic than gossypol and melts at about 246-248°. Again, by fusion with alkalı, a second new crystalline substance. C-gassypol, can be obtained, and this, which is soluble in alkalis with a blue colour, is probably elosely releted to the purple oxidation product of gossypol referred to above.

D.Gossypol,-Cotton seed meal which has been submitted to the "cooking process" loses much of its toxic property, and then contains practically no gossypol. By extracting this product with aniline, the aniline compound of D-gossypol can be isolated (yield, 12%) Alcoholie potsah converts this substance into D-gossypol, yellow crystals which darken and soften at about 256° (Carruth, Ic.),

Sherwood (J. Agric, Res. 1926, 32, 793) seed meals 75% or more of the gossypol present is converted into the less toxic D gossypol, and this he suggests is a hydrolysis product of the unstable gossypol According to Gallup (Ind Eng. Chem 1927, 20, 59), the extent to to be an a hydroxy-acid Methyl methylogony (Ind Eng. Chem 1927, 20, 50), the states to polate, [C₁;H₁,O₂](OHe)₁la, mp. 142°, gives which gosypol is converted into D gosypol methylogosypolic acid, mp. 225°, by hydrolysis (Karrer and Tobler, 1-c). especially in the presence of water, destroys the D-gossypol, finally giving a product free from both forms of gossypol and almost non toxic

According to Clark (l.c.), D gossypol yields

diaminogossypol with aniline. A. O, P, and E. J. C. COTTONSEED OIL, an important by product of the cotton industry is prepared on an enormone scale from the seeds of the several species of the cotton plant, Gossypium sp , which growe wild as a perennial shrub in the tropics, and is extensively cultivated (ss an annual crop) for the cake of the fibre in hot but non-tropical regions in many parts of the world In the United States the "Upland cotton," G Airsutum L , in which the seed is covered with a short fuzzy down (" linters ") in addition to the longer cotton fibre, is cultivated, whereas in the Sea Islands and Egypt, G. barbadense is grown, which furnishes "naked" or "bald" seeds, from which all the cotton fibre (but) can readily be etripped. The Indian and Asiatic venetics cotton include the ill-defined epecies G. herbaceum, G. arboreum, G. indicum and G. neglectum, whilst the indigenous South American cotton plants are frequently classified carbonyl groups originally present. opoGossypol as separate species G. acuminatum (=G. peru-is considerably less toxic than gossypol. It wanum Cav.) and G. brasiliense Mecf. (=G. perumanum Engl.). Whilst the native cottons yield almost naked seeds, fuzzy seeds from cultivated varieties of "Upland" cotton are also exported from Brazil.

For classification of the cottons, are Watt,

No. 12, 1935; T. H. Kearney and G. J. Harrison, J. Agric. Res. 1928, 37, No. 8, 465; H. B. Brown, "Cotton," 1927; Angelasto, Boyle, Meloy and Stine, "The Cotton Situation," U.S. Dept. Agric. Year Book 1921, 323).

In this country, 1 cottonseed appears to have been first expressed in 1847, and refined edible cottonseed oil was being produced in France, from Egyptian seed, in the middle of the nineteenth century. In the United States, however, the development of the industry, hampered as it was by the lack of proper methods for decorticating the seed and for refining the oil, and further checked by the Civil War, was very slow; prior to 1860 practically the whole of the seed not required for planting was wasted, and although numerous attempts at oil production were made from 1820 onwards, the commercial establishment of the cottonseed industry in America and its rapid progress may be said to date only from as late as 1867-1870. (For the early history of the cottonseed industry, cf. Lamborn, "Cotton Seed Products," New York, 1904.) Since then, and especially within the present century, the industry has expanded to such enormous dimensions that the United States. which formerly exported large quantities of cottonseed oil to Great Britain and Europe, now not only consumes internally the whole of her vast crop of over 3-4 million tons p.a. of seed (equivalent to about 510,000-700,000 tons of oil), but imports in addition a considerable amount of cottonseed oil (57,000 and 86,600 tons in 1936 and 1937 respectively) from the United Kingdom (from one-third to one-fifth of the total), Japan, Brazil and Egypt. The annual import of cottonseed into the United Kingdom (chiefly from Egypt, Sudan and British East Africa, and Brazil) amounts to some 600,000-700,000 tons, and the export of oil to some 23,000-40,000 tons. India, which was formerly an important supplier of seed to this country, has ceased to export seed since about 1932, the whole crop (c. 2,000,000 tons p.a.) being consumed locally and only the expressed cake being exported. Large quantities of cottonseed are also grown (chiefly for local use) in South Russia (c. 850,000 tons p.a. from 1931-1934) and China (c. 1,250,000 tons p.a.).

About 1,000 lb. of seed are obtained per bale (500 lb.) of cotton fibre produced. The seed consists of about 40-55% of shell ("hull") and 45-60% of kernel, and contains as a rule from 6 to 12% of water, 15 to 25% of oil, 19 to 24% of proteins, 24 to 31% of carbohydrates, 14 to 21% of crude fibre, and 3 to 4.5% of ash. The proportion of oil varies with the origin of the seed as well as with local climatic conditions; thus on the average, whilst Egyptian seed contains up to 24% of oil and South American seed from 20 to 21%, East Indian seed averages about 18% and American seed about 19.5% of oil. For commercial statistics, the oil equivalent

of cottonseed is taken as 17 or 18%.

A distinction must be made in practice between the "naked" or "bald" seed from

Egypt, etc., which is processed in England, France, etc., and can be crushed and expressed directly, and the seed from the "fuzzy" varieties of cotton which need to be decorticated before expression, since, even after ginning and delinting, the seed-hulls retain a fuzzy coating of short fibres, the presence of which in the seed-cake would be undesirable. After removing as much as possible of this "linters" (which is used in the manufacture of guncotton, paper and other cellulose derivatives) in gin-like "delinting" machines, the seeds are passed through "hullers" in order to crack the hulls, which can then be separated from the kernels or "meats" by sifting through shaker screens. Chemical methods of removing the linters, e.g. by treatment with acid fumes (see Ardashev, Ind. Eng. Chem. 1933, 25, 575; Ardashev and others, Cotton Ind. Trust, U.S.S.R., Coll. Papers, 1933, No. 1, 38, 40, 55, 83), are less frequently resorted to except for analytical purposes. The "meats," or in the case of the "bald" varieties the whole seeds, are then crushed and "cooked" (i.e. heated in the presence of open steam) before expressing the oil in hydraulic presses. Both the Anglo-American and the cagetype of press are commonly employed, bnt, of recent years, the continuous acting expeller type of press has also been employed in the United States; such "expeller oil" is still frequently denoted by the misleading term " cold-pressed oil," although, in fact, the crushed seed is nowadays steamed and pre-heated before being passed into the expeller.

The extraction of the oil from the seed by means of solvents has also been proposed (see D. Wesson, Oil and Fat Ind. 1933, 10, 151; de

Raedt, Mat. Grasses, 1935, 10673).

Any "foots" or fragments of crushed seedmeal are separated or filtered off from the oil before storage for sale as "crude cottonseed oil." The separated "hulls" may be used as bedding for cattle, as a fertiliser, or as fuel; according to Jamieson ("Vegetable Fats and Oils," A.C.S. Monograph, 1932; cf. Kress, Cotton Oil Press, 1930, 3, No. 12, 33) increasing quantities, after removal of the "hull-fibre" (linters) in special machines, are ground to make 'hull-bran,' which is added to the expressed seed-cake intended for use as cattle-food. Proposals have also been made to work up the hulls by destructive distillation (see Randolph, Grove and Tucker, J. Elisha Mitchell Sci. Soc. 1932, 48, 26) or for the recovery of pentosans, furfural, etc. (see Zakoshikov, Ivanova and Kurennova, Cotton Ind. Trust, U.S.S.R., Coll. Papers, 1933, No. 1, 87; Zukervanik and others, Bull. Univ. Asie Centr. 1935, No. 21, 69; Kao and Yü, J. Chem. Eng. China, 1936, 3.

Expressed cottonseed cake (or meal) serves as a valuable ingredient of cattle-food, whilst inferior grades are employed as fertiliser; the use of cottonseed meal for human consumption has been suggested (see Wesson, Oil and Fat Ind. 1926, 3, 121; Baumgarten, Cottonseed Oil Mag. 1924, 40, No. 10, 28).

Standard methods for the analysis and evaluation of cottonseed, including determinations of oil, nitrogen, etc., have been published by the

¹ As early as 1783-1789 a prize was offered by the Society of Arts of London for a satisfactory process for the preparation of oil and cattle-cake from cotton-seed, but this was never claimed.

Amer. Oil Chem. Assoc. (Standard Methods of | "Extracted" and "Cold-Pressed" (1 c. ex-Analysis); U.S. Dept. Agric., Service and Reg. poller-) oils are recognised (see "Book of Rules"
Announcements, Bur. Agr. Economics, No. 133, of the National Cottonseed Products Assoc 1932 : Assoc. of Off. Agric. Chem. (Official and Standard methods for the axamination of crude Tentative Methods of Analysis); see Jamieson cottonseed oils, including determination of and McKinney, Oil and Fat Ind. 1930, 7, 291; Meloy, 18td. 135; 1931, 8, 187; for the deter-mination of "lint" and "bull fibre," see also Official Methods of the Amer. Oil Chemists' Freyer, Oil and Soap, 1935, 12, 208, 259; McKinney and Jamieson, ibid. 1936, 13, 139.

cells containing a deep red brown colouring matter, with which is associated a highly toxic phenolic substance "gossypol," C₃₀H₃₀O₈ (see COTTON SEED, Colouring Matter of)

Some of this toxic material, which occurs to the extent of about 0 4-1-2% of the kernel (see ("soap atock") falls to the bottom of the Schwartze and Alsberg, J. Agrae. Res. 1923, 25, 285; Gallup, Oil & Soap, 1936, 13, 191), may pass into the oil, if this is really expressed in the cold ; however, under the action of moist heat, incidental to the "cooking" process in commercial oil recovery, the gossypol appears to comane with the seed proteins and is converted a declared as required. Oils intended for into a harmless substance known as "house declared purposes may be treated with chemical goaypol," or "D goaypol," or that the declared purposes may be treated with chemical and intended for leading agents.

The best qualities of refuned oil, which have locally 1, Agrin (Res. 1923, 6, 321 ; 1923, 6, 30. A, 15. Merand, 1323, 5, 333. N.B.—The difference occasionally locally representations of the control of the combine with the seed proteins and is converted observed to aruse from the over feeding of cattle on cottonseed meal, which were formerly attributed to the action of gosaypol, are now conaidered to he, in large part at least, due to the lack of eertain proteins and other dietery factors in the meal, and can be remedied by the addition of appropriate food supplementa manufacture of various types of soap and soap to the feed (see Clark, i.e.; Gallup and Reder, powders. The Indian and Egyptian cottonsed J. Agric, Res. 1935, 51, 259; 1936, 52, 65); for the determination of total and "hound" gossypol in cottonseed meal, see J. O. Halverson and Smith, Ind. Eng. Chem. [Anal], 1933, 5, 319; 1934, 6, 356; 1937, 9, 516.

Correspondingly, only traces of the order of 0.05% of gossypol (? "bound" gossypol) are to be found in commercial hot-pressed crude cottonseed oil see Royce, Oil and Soap, 1933, 10, 183; 1934, 11, 116); larger amounts (up to 1%) have been found in expeller-pressed and in solvent-extracted oils. However, any gosaypol present in the crude oil is completely removed during the normal refining treatment with

alkalı. Owing to the presence of the colouring matters (which include plant pigments and resins, see Jamieson, Oil and Fat Ind. 1925, 2, 101; Thornton, jun, ibid. 1934, 11, 209), erude cotton seed oil has a ruhy red to almost black colour, the depth of the colour depending in the first off the deposited glycerides; the filtered oil, instance upon the freshness of the seed. Seed known in commerce as "uniter oil" "dearn that has been stored for a long period may under garinated" o'", it is required go deterioration ("heating"), which is reflected in the interior guilty of the all obtained the

" refining loss" and the behaviour in bleaching Society. The usual determination of "refining loss" has been modified by McLeod (Oil and Raw cottonseed kernels are studded with Soap, 1936, 13, 67) in order to render it applicable

The crude oil is refined by treatment with dilute caustic soda at a temperature not ex ceeding 120°F. (49°C.), whereby the frea fatty acids are neutralised, and the soap formed vessel, carrying with it the brown colouring matter and other impurities, so that the supernatant nentral oil is of a bright yellow colour This oil, which is washed to remova traces of ecap, is known as "summer yellow oil," and may be bleached with adsorbent earths and deedorised as required. Oils intended for

16% is employed as a salad or cooking oil, 2% for margarine manufacture, and 70% (much of it after hardening by catalytic hydrogenation) in the manufacture of shortenings (lard subattutes); the halance of 12% of technical oil, and also the fatty acids recovered by working up the "aoap stock," are chiefly utilised in the

oils ere used to a greater extent in soap-making By the destructive distillation of cottonseed oil combustible gases and hydrocarbon oils tan bo produced (see Egloff, Ind. Eng Chem 1932, 24, 1426; Martin, Bull, Mat. Grasses, 1932, 16, 97), and it is possible that the application of auch processing to low-grade crude oils, or to cottonseed itself, may have a certain commercial aignificance in remote districts, such as Central Africa, where imported petroleum products are

abnormally expensive. Refined cottonseed oil (" summer oil") is, at ordinary temperatures, a pale yellow oil of characteristic but not unpleasant flavour; on atanding at about 12°C, a deposit of "steam, s e. higher melting glycerides, is formed. As this "atearm" would be objectionable in the case of oils intended for use as table or salad oils, it is removed on the larga scale by chilling the oil to a low temperature (5° C, or lower) and filtering go deterioration in decising A which is therefore the first property of the form Croid actionsed oil is valued and from Croid actionsed oil is valued and according to its flavour, acutety, refaming loss, and the quality of the refined oil producing. The specific gravity of refined from it, in the American trade, 6 grades from 0 925 at 15 5°C, ; the saponification rather varieties. The refined is the control of from 100 to 100 and the follower value from 100 to 100 to 100 and the follower value from 100 to 100 and the fo

present. On the whole, the oil from Egyptian seed tends to have a lower density and iodine value than the North American product (see Jamieson and Baughman, J. Amer. Chem. Soc. 1920, 42, 1197; Oil and Fat. Ind. 1926, 3, 347; 1927, 4, 131; Perdrigeat, J. Pharm. Chim. 1930, [viii], 12, 307); thiocyanogen values of 63-64.8 are recorded for three Egyptian oils hy Atkinson (Analyst, 1934, 59, 399). Analyses of cottonseed oils by Jamieson and Baughman (l.c. 1927) and Hilditch (Hilditch and Rhead, J.S.C.I. 1932, 51, 200T; Hilditch and Jones, ibid. 202) show that the fatty acids of cottonseed oil contain from 19 to 22% of palmitic acid, 24 to 34% of oleic acid, and 39 to 50% of linoleic acid; only from 1.5 to 2.5% of stearic acid is present together with small amounts (0.5–3%) of myristic acid (?) and arachidic acid (<1%). Practically the whole of the palmitic acid is associated with unsaturated fatty acids in mixed glycerides, only about 1.5% of fully saturated glycerides heing present; the balance of the oil appears to consist of about 60% of mono-palmito-glycerides, 15% of di-palmito-glycerides, and 24% of mixed triglycerides of oleic and linolic acids (Hilditch and Jones, J.S.C.I. 1934, 53, 13T). S. Ueno and Iwai (J. Soc. Chem. Ind. Japan, 1935, 38, Suppl. 602) report the following composition for the fatty acids in a sample of fully hydrogenated cottonseed oil: myristic acid, trace: palmitic acid,

17%; stearic acid, 76%; higher saturated acids (arachidic acid), 7%.

Besides the sterols (unsaponifiable matter, which includes β -sitosterol, stigmastanol and waxy matter, see E. S. Wallis and Chakravorty, J. Org. Chem. 1937, 2, 335; Anderson and Moore, J. Amer. Chem. Soc. 1933, 45, 1944), traces of plant pigments (flavone derivatives), and phospholipins are also present in the refined oil; these occur in greater amount in crude cottonseed oil, together with a number of other minor components—raffinose, pentosans, pro-teoses, peptones, phytosterolins, inosite phosphates, etc. (Jamieson and Baughman, J. Oil and Fat. Ind. 1924, 1, 31; 1925, 2, 101; 1926, 3, 153, 347); a small hut bulky precipitate (known as "settlings") which occasionally settles out from a filtered crude oil consists of a mixture of such adventitious "im-

purities."

Cottonseed oil typifies a semi-drying oil: in the Livache test it absorbs 5.9% of oxygen in 24 hours. By blowing air through cottonseed oil at a temperature of 90-100°C., oxygen is absorbed with the formation of a small amount

to 115; ahout 1% of unsaponifiable matter is | Halphen colour test 1 is most useful, this test is usually carried out as follows: 1-3 c.c. of the oil is dissolved in an equal volume of amyl alcohol; 1-3 c.c. of a 1% solution of flowers of sulphur in carhon disulphide is added, and the test-tube is immersed in boiling water (or, better, boiling brine); the heating is continued for 2 hours or until a red coloration, indicating the presence of cottonsecd oil, appears (usually within 20-30 minutes); the intensity of the colour depends in the first instance upon the proportion of cottonseed oil in the sample, but also varies somewhat with different specimens of this oil. The reagent which produces the colour appears to be an impurity in the amyl alcohol (Gastaldi), and the test may he rendered more delicate by replacing this solvent by pyridine; in this way, it is claimed that 0.25% of cottonsecd oil may he detected (Gastaldi, Annali Chim. Appl. 1914, 2, 203; Pieraerts and Simar, Mat. Grasses, 1928, 8222, 8312; see "German Standard Methods for the Analysis of Oils and Fats "(Wizoeff). According to Shelley (Analyst, 1925, 50, 132; see Gastaldi, l.c.; "British Pharmacopœia," 1932) hetter results are obtained with the original Halphen reaction if the testtuhe is sealed before heating. Although the Halphen test cannot be used for quantitative analysis, it must be considered, if positive, as one of the best means for the qualitative detection of cottonseed oil; a negative result, however, is no conclusive proof of the absence of this oil in a mixture, as the colour-producing constituent may he destroyed hy heating or blowing the oil, or even hy rancidification on prolonged storage; the test may also fail with hydrogenated cottonseed oils.

COUCH GRASS. Associated in couch grass with triticin (a carhohydrate resembling inulin) are dextrose, mucilage, mannitol, and inositol) (v. Agropyrum).

COUMARANE. Coumarane, or hydrocoumarone, is prepared by reducing coumarone in alcoholic solution with sodium (Alexander, Ber. 1892, 25, 2409). It forms a colourless oil, b.p. 188-189°, and gives in sulphuric acid hy addition of ferric chloride a violet coloration.

absorbed with the formation of a small amount, of "oxidised acids," while the acetyl value, viscosity, and density rise considerably. This reaction can be applied on the large scale to the preparation of blown cottonseed oil which was formerly used to a considerable extent in admixture with mineral oil to form lubricating oils of the marine oil type.

As stated above, hydrogenated (hardened) cottonseed oil is used to an enormous extent as an ingredient of shortenings.

For the identification of cottonseed oil, and for the detection of it in other oils and fats (down to 1% in favourable circumstances), the

COUMARIN.

was obtained by Vogel (Gilb, Ann. 1829, 64, 163) from the tonks bean, from mehlot, and from woodruff but was mistaken for benzoic acid. It was first recognised as a distinct anbstance by Guibort, who gave it the name coumarin. It is a normal constituent of lavender oil and is very widely distributed in the plant langdom (Gildemeister and Hoffman, "Die atherischen Ole," 1928, Vol. I., p. 652). It occurs in plants as a glicoside which is hydrolysed by an enzyme present in the plant, for its determination in the plant. planta, see Duncan and Dustman, Ind Eng. Chem. [Anal], 1934, 6, 210.

It is prepared from tonka beans with 80% alcohol, followed by precipitation with water with a small amount of 8 nitrocommarin (De. (Wohler, Annalen, 1856, 98, 66); it may also and Krishnamurthi, J. Indian Chem. Soc. 1927. conveniently be extracted from melilot (Ober-

mayer, Z. anal, Chem 1913, 52, 185)

It was first prepared synthetically by Perkin (Perkin, J.C S. 1871, 24, 52; Sep and Chakra by heating the sodio derivative of salicylaldehyde with acetic anhydride (JCS, 1868, 21, 53) or, better, by boiling together salicylaldehyde, studsed by a number of workers (Sen and acetic anhydride and anhydrous sodium acetate Chakravarti, 161d. 1929, 8, 847; Naik and Patel. (idem, ibid 1887, 51, 388, see Tiemann and JCS 1934, 1043; Chakravarti, J. Indian Chem Herzfold, Ber. 1877, 10, 284). Yanagasawa and Soc 1935, 12, 139) and the introduction of Kondô (J. Pharm Soc Japan, 1921, 498) have described an improved method, using sodine as and Das Gupta (1bid. 1931, 8, 417). a catalyst, which is claimed to give a 70% yield Perkin's method is that most frequently used for the technical production of coumarin. A variant of the process consists in the treatment of salicylaldehyde with acetic anhydride at 180°

(Reychler, Bull, Soc chim, I897, [m], 17, 516).
Another method used commercially is the condensation of salicylaidshyde with malonie acid in acetic acid, followed by deearboxylation

hy heating the coumann 3 carboxylie acid so

formed (Stuar, J.C.S. 1886, 49, 267).

Coumann is also formed by hesting a matture of potential and a student of potential and a tended to the synthesis of substituted commarine, phenol homologues being employed (see Bailey and Boettner, Ind Eng Chem. 1921, 13, 905) The most important method of ohtaming substituted coumarins is by the condensation of mination in vanilla essence, see Winton and phenols with ethyl acetoacetate or its alkyl Silverman, J. Amer. Chem. Soc. 1902, 24, 1123: derivatives in the presence of sulphuric acid; Winton and Bailey, abid. 1905, 27, 719, thus, phenol and cthyl acetoacetate yield 4 methylcoumarin (von Pechmann and Duisberg, Ber. 1883, 16, 2127; von Pechmann and von Krafft, ibid. 1901, 34, 421, Peters and Simones, ibid. 1908, 41, 831). Phenois containing alkyl, hydroxyl, or dialkylammo groups in the m- or 2:4. positions give good yields in the reaction, 2:4- positions give good yields in the reaction, but condensation is prevented by the presence of initio, carbox 1 or carbothoxy groups (Layton, J.C.S. 1903, §2.018). Phosphore acid is said to be superior to sulphurio acid as a catalyst in for this reaction (Chartwarti, J. Indian Chem. Soc. 1931, 8, 613; 1935, 12, 253) while city achieves the superior of the property of the superior of the super

densation in good yield when the ordinary method fails (Appel, J.C.S. 1935, 1931).

Coumarin forms rhombic crystals, mp. 70. h p. 291°. It has a pleasant aromatic odors and is aparingly soluble in hot water, readily soluble in boiling water, and very soluble in alcohol and ether. Crude coumarin may be purified by repeated crystallisation from light petroleum (Claassen, Pharm. J. 1897, 4, 181) Hot concentrated alkalis convert comman

into salts of coumario acid (case hydrory cinnamic acid) from which coumarin is re formed hy heating with acetic anhydride. Reduction of coumarin with sodium amalgam in water or alcohol or with sinc and addium hydroride yields mehlotic acid (Zwenger, Annalen Suppl. 1867, 5, 122; 1872, 8, 32; Fries and Fickewith Annalen, 1908, 362, 30, 35). On nitration 6 nstrocoumarin is the principal product (Dela lande, abid 1842, 45, 337; Morgan and Micklethwait, J.CS. 1904, 85, 1233) together with a small amount of 8 mitrocommann (De) 4, 197). Suiphonation yields, first, coumant 6 sulphome acid and then the 3:6 disulphomic sed varts, J. Indian Chem. Soc. 1928, 5, 433)

The mercuration of commanns has been arsenic into the coumsrin nucleus by Goswami

Sodium bisulphite reacts with coumarins to give \$\beta\$ aulphonic acids (Dodge, J. Amer. Chem Soc. 1916, 28, 448; 1930, 52, 1724; Dey and Row, J.C.S. 1924, 125, 554), a reaction that has successfully heen applied for the determination of commarin (Radchife and Sharples, Perf. and Essent. Oil Rec. 1924, 15, 396, 437; 1925, 16,

20 et seg). In Germany, woodruff is steeped in white wine, to which it imparts the pleasant groms of

Contractor, but the agreeable odour of superila and is used in perfumery for the preparation of asperula essence. It is also frequently used to adulterate vanilla extract. For its deter

H. N. R. Coumarm COUMARIN GLYCOSIDES.

E. F. A.

sides are:

. 7-hydroxycoumarin. SkimminÆsculin . 6:7-dihydroxycoumarin. . 6:7-dihydroxycoumarin. Cichorin Daphnin . 7:8-dihydroxycoumarin. . 6-methylæsculetin. Scopolin . 6-methylæsculetin. Fabiatrin . 6-methoxy-7:8-dihydroxy-Frazin . coumarin.

Skimmin has the glucose attached at position 7, as also has daphnin. In æsculin it is at r, as also has daphine. In second it is at position 6 (Head and Robertson, J.C.S. 1930, 2434), in cichorin at position 7 (Merz. Arch. Pharm. 1932, 270, 476), and in fraxin at 8 (Wessely and Demmer, Ber. 1928, 61, [B], 1279; 1929, 62, [B], 120). Scopolin contains the charge perioder of historic archaeographics. two glucose residues, fabiatrin only one.

COUMARONE, 2:3-benzfuran,

occurs in the coal-tar fraction of b.p. 165-175°. After removing pyridine bases and phenols by treatment with acid and alkali, the oil is treated with bromine at 0°, and the dibromide thus obtained reduced to coumarone with alcoholic potash and sodium amalgam (Kraemer and Spilker, Ber. 1890, 23, 78; Burda and Sukat-scheva, Ukrain. Chem. J. 1931, 6, [Sci.], 169). It may also be obtained from this source by adding picric acid to the oil, when coumarone picrate crystallises out. This is separated, decomposed with hot water or dilute alkali, and the coumarone obtained by steam distillation. The picric acid is recovered and used for another operation (G.P. 53792; B.P. 1422, 1890). It may be separated from accompanying indene by condensation of this with ethyl oxalate (Thiele, Ber. 1900, 33, 3400).

It is obtained synthetically by heating o-hydroxy-ω-chlorostyrene with caustic potash (Komppa, Ber. 1893, 26, 2971), by heating phenoxyacetaldehyde with zinc chloride in acetic acid (Stoermer, Annalen, 1900, 312, 261), and by heating phenoxyacetaldehyde diethylacetal with anhydrous oxalic acid (Stoermer, l.c.).

Coumarone is a colourless oil of aromatic odour, b.p. 173-175°/760 mm., 62-63°/15 mm., $d_4^{22\cdot7}$ 1.0913, $n_D^{22\cdot7}$ 1.5645, and is insoluble in water but soluble in alcohol and ether. It may be characterised by means of its picrate, m.p. 102-103°, and 1:3:5-trinitrobenzene m.p. 102-103°, and 1:3:5-trinitrobenzene derivative, m.p. 104°. It is stable to alkalis, ammonia, potassium cyanide and hydrochloric acid but yields a series of polymers with sulphuric acid (v. Coumarone Resins). The coaltar fraction, b.p. 185-195°, contains methylcoumarones (Stoermer and Boes, Ber. 1900, 33, 3013).

H. N. R. COUMARONE RESINS. Kraemer and Spilker (Ber. 1900, 33, 2257; 1901, 34, 1887) either green or dry, and t found that coumarone was polymerised by

The aglucones of the hydroxy coumarin gluco-| treatment with sulphuric acid; Heusler (Z. angew. Chem. 1896, 9, 318) found that aluminium chloride had a similar action, and Stein (ibid. 1919, 32, 246) showed that stannic chloride was also an efficient catalyst for this polymerisation.

Technical coumarone resins are mixtures of coumarone and indene polymers obtained by treatment of coal-tar naphtha of b.p. 150-200° with suitable catalysts, notably sulphuric acid. Ellis and Rabinovitz (Ind. Eng. Chem. 1916, 8, 797) made a thorough study of this process and concluded that sulphuric acid was the most effective catalyst; ferric chloride or aluminium chloride (G.P. 446707) and fuller's earth mixed with ferric sulphate (U.S.P. 1894934) have also been claimed to be effective catalysts, while according to Bojanowski, Giziński and Rabek (Przemysl Chem. 1934, 18, 321) almost colourless resins are obtained by using a mixture of sulphuric acid, acetic acid and water in the ratio 15:4:1. For discussions of the technical procedure, see Marcusson, Chem.-Ztg. 1919, 43, 93, and Glaser, Brennstoff-Chem. 1921, 2, 99, 113.

The best quality coumarone resins are pale amber solids melting above 130°. They are classified according to colour and hardness; resins softening above 50° are termed "very hard," between 40° and 50° "hard," and between 30° and 40° "medium hard." For methods for their examination, see Marcusson, Chem.-Ztg. 1919, 43, 109. Coumarone resins differ from many other resins in that they are neutral and unsaponifiable; although somewhat unsaturated they are unaffected by most reagents; on heating above 300° they undergo partial depolymerisation. Technically, they find wide application, being especially useful in the manufacture of varmishes, printing inks, waterproofing materials, rubber compositions and chewing gum. For a complete review of the chemistry and application of coumarone resins, see Ellis, "The Chemistry of Synthetic Resins," 1935, Vol. I, ch. 5-7.

H. N. R. COVELLINE or COVELLITE. Native cupric sulphide, CuS, crystallising in the hexagonal system. Crystals are rare and have the form of thin six-sided plates. Fine large crystals have been found at Butte in Montana. The mineral has a characteristic indigo-blue colour, hence the name indigo copper (Ger. Kupferindigo). It occurs in most copper-mines as a blue, earthy coating on other sulphide ores of copper, and is sometimes found in sufficient quantity to be of importance as an ore, e.g. at Butte in Montana, and Copiapo in Chile. Covellite containing platinum (as the mineral sperrylite, PtAs₂) is mined in the Medicine Bow Mountains in Wyoming.

COW PEA or CHINA BEAN. The seed of the legume Vigna sinensis (L.) Endl. (or Dolichos sinensis, L.), are variously coloured, approximately ½ in. long, and are formed in long narrow pods often more than a foot in length. The peas are eaten as a vegetable, either green or dry, and the whole plant serves

Analyses of the besns average :

			Edible	US Dept.	Ebodesian Dept.
			beans.	(1906).	Agric, (1934)
			.%	%	%
Water			66 0	148	139
Protein			94	208	23 4
Fat			06	1.4	1.8
N.free e:	rtra	ct.	22 7	65 7	59
Fibre .			_	4-1	51-6
Ash			7.4	3.9	3.4

According to Osboroe and Campbell (J. Amer. Chem. Soc. 1897, 19, 494) the protein constituents of the cow pea include 3 globulines (i) Vignin, the major constituent (C 52 6, N 173, S 0 8%); (u) phaseoin (C 52 3, N 167, S 0 5%); and (ui) a soluble globulin (C 53 2, N 164, S 11%).

Oaborne and Heyl (Amer. J. Physiol. 1908, 22,

414

362) found the products of hydrolysis of signin

	%			%
Glycocoll	0.00	Phonylalanın	0	4 6
Alanune	0.78	Probine		4.4
Valine .	0 29	Tryptophan .		+
Lencine	6 73	Arginine		60
Aspartic acid	. 294	Lysine		37
Glutamic acid	. 1284	Histidine .		27
Tyrosine .	2 03	Ammonia .		21
Jones et al (J. 1 rignin to contair				

1.65%. Adolph and Chiang (Chinese J. Physiol.

1935, 9, 347) separate five fractions among the proteins of eow pes, viz.

Vignin (Glob	nlu	a A)		. 45
Glutelin .					. 25
Albumin					15
Globulin B					. 10
Globulin C	٠				. 5

•

and record the high cystine content of the globulins and albumin and the high lyaine content of the glutelin. Bhagvat (J. Indian Inst. Sci. 1935, 18, A, 39, 137) also examined the proteins and confirms the high cystine content of the globulus. Averago analyses of the green parts of the

cow pea as used for fodder are recorded by the U.S. Dept. of Agriculture as :

Creen folder

					٠	100	i roduer,	24 %
Water							83 6	107
Protein		٠					24	166
Fat .		٠.				٠	04	29
N-free e	ztr	act					7-1	42.2
Fibre.	•	•	•	٠	•	•	48	20 1
Ash .	•		•		٠	•	1.7	75

A. G. Po "CRAB'S EYE " v. ABRUS PRECATORIUS

CRANBERRY. The fruit of the American bog plant Vaccinium macrocarpon Ait. A amalier apecies (V. Ozycoccus, L.) occurs both in Europe and America. A related species (V. ertio-idata) is known as the mountain or red eranberry, red whortleberry or cowberry in 11 3, protein 0 12, acida (as citric) 1-9, invert England, and as forherry in the United States. sugar 6 5, sucrose 0 5, tannin 0 19, ash 0 3%. The fruits are notable in keeping for a long time | Vaccimin is also present in the fruit.

without decomposition, largely because they contain appreciable amounts of benzoic and The extremely sour fruit find widespread use in the form of jam or jelly.

Average analyses of cranberries are :

	American,	European.
Water	87-8-88 8	89 3–89 6
Protein .	0 35-0 5	0 10-0-12
Fat	0 57-0 97	_
N. free extract	9 0-10 1	101
Free acids 1 .	23-24	23
Sugars	3 5-4-1	1.5
Fibre	1-1-1-5	_
Ash	0 15-0 18	0 15-0 17

1 Calculated as citric sold,

Cranberry juice contains invert augar 87 L acids (calculated as malic scid) 190, ash 30, netrogen 0 11 g. per litre, and either because of the benzore acid content or because of the small amount of nitrogen present is not fermentable by

While all investigators agree on the presence of benzoic acids (0 02-0 09%) in cranberries, there seems some difference of opinion as to the nature and proportion of other acids occurring in the fruit Citric and malic acids are generally reported in American cranberries ; Stolle (Z. Ver deut Zucker Ind. 1900, 27, 609) found glyoxyle acd in European fruit, and Rising (Kgl Landtbruks Akad. handl. Tidakr. 1914, 329) also found reovalorie acid. According to Isham et al (Massachusetts Agric. Exp. Sta Bull. 1935, No. 315) make acid is present in mature and mmature fruit in similar proportions (0 26%), whereas the citric acid content declines some what (1.31-107) with advancing maturity The same authors also record more than 0 5% of quinic seid in the fruit.

Rising (l c.) also isolated from cranbernes the glucoside vacciniin.

CaH,,Oa COCaHs.

The ash of American cranberries contains: K₂O 480, Na₂O 66, CaO 186, MgO 68, Fe₂O₃O 7, P₂O₃143, SiO₂ 622, Cu 0009%. According to Willstätter (Sitzungsber, K. Akad. Wiss. 1914, 12, 402) the pigment of the cranberry is an anthocyanin (chlonds, C15H11OcCI), a compound of galactose with cyanidin. Claassen (101d. 16, 70) isolated from the fruit a bitter substance ozycoccia which

gave reactions similar to those of arbutin. The cowberry or foxberry closely resembles cranberry in nature and uses Recorded analyses by Ystgaard (Tids. Norske Landwbraks. 1902. 9, 125) give the average composition as: total solids 155, protein 0.25, total acids 1.92, citric acid 054, malio acid 031, sugars 64, pentosans 059, fibre 185%. According to Hotter (L landw Veruschew, 1909, 9, 147) the segars comprise glucose 3 0-4 8, fructose 40-58, sucrose 04-08% of the total dry matter. Windisch and Schmidt (Z. Unters, Nahr.

Genussm. 1909, 17, 584) show the average composition of the juico (sp gr. 1-043) to be; solids

1912. 24, 575) give the following analysis:

Water. % 5-97	Protein. %	Fat. %	N-free extract and fibre. % 38-56	Ash. % 2·11
0.91	40.44	30.12	30.00	~ 11

The seeds yield a drying oil having the following characteristics: d_{15} 0.9301, refractive index decomposition of creatinine zinc chloride by 1.4753 (25°), saponification value 190.1, iodino value 169.2, Reichert-Meissl value 0.55, Polenske value 0.30, Hehner number 95.7, acid value 1.94, hexabromides of linoleio and linolenic acids 22.8%.

A. G. Po.

CREAM OF TARTAR. Acid potassium tartrate, C₄H₅O₆K.
CREAMING v. BLEACHING.

CREATINE. Methylglycocyamine, Methylguanidinoacetic acid,

H₂N·C(NH)NMe·CH₂·CO₂H,H₂O,

discovered by Chevreul (Annalen, 1832, 4, 293) in muscle (xpéas=flesh), occurs in human flesh, in the flesh of many warm and cold blooded animals, and in urine under certain conditions (von Liebig, ibid. 1847, 62, 257; Gregory, ibid. 1848, 64, 105; Schlossberger, ibid. 1848, 66, 80; Price, ibid. 1850, 76, 362; Voit, Z. Biol. 1868, 4, 77; Marcet, J.C.S. 1864, 2, 406; Folin and Denis, J. Biol. Chem. 1914, 17, 488; Rose, ibid. 1911, 10, 265); the amount varies with the species and the organ; the flesh of fowl yields 3.21%; of pigeon, 0.825%; of ox, 0.697%; of ox heart, 1.375%; of herring, 1.324%; and of brill, 0.614% (Gregory, l.c.; Beker, Z. physiol. Chem. 1913, 87, 28; according to Myers and Fine (J. Biol. Chem. 1913, 14, 9) it is constant for a given animal.

The origin and physiological significance of creatine and creatinine are still controversial subjects of too detailed a nature to be considered here. Creatine is very unevenly distributed in the body, about 98% being found in the muscles and most of the remainder in the brain. It is not found in the urine of normal adults and cannot be regarded as a waste product of the metabolism of protein from which it is presumably formed. It is an essential substance for muscle metabolism since a large part of the creatine in muscle is present in combination with phosphoric acid as phosphagen (phosphocreatine); the hydrolysis and resynthesis of this compound are important phases in the processes of muscular contraction and relaxation, phosphagen thus serving as a reservoir for the inorganic phosphate necessary for the process of contraction. It is probable that creatinine is a waste product formed from creatine, although this has been disputed. For further details and refer-ences, see Hunter "Creatine and Creatinine," Longmans, Green & Co., 1928, Peters and van " Quantitative Slyke Clinical Chemistry." Baillière, Tindall & Cox, 1931, and Lovatt Evans "Recent Advances in Physiology," Churchill, 1930.

Cowberry seeds, according to Diedrichs (ibid. | trate is treated with basic lead acetate until no further precipitation occurs, and again filtered after removing the excess of lead from the filtrate by sulphuretted hydrogen, when it is evaporated until creatine separates on cooling; this is washed with alcohol and recrystallised from water (Neubauer, Z. anal. Chcm. 1863, 2, 26; 1867, 6, 33). Creatine may be prepared by the means of calcium hydroxide (Benedict, J. Biol. Chem. 1914, 18, 186).

Creatine is prepared synthetically by heating sarcosine with cyanamide at 100°:

CN·NH₂+NHMe·CH₂·CO₂H =NH:C(NH2)NMe·CH2·CO2H

Volhard, Z. Chem. N. F. 1869, 5, 318; Strecker, Jahresber. Chem. 1868, 686); or, together with creatinine, by fusing sarcosine with guanidine carbonate (Paulmann, Arch. Pharm. 1894, 232, 638). It has also been prepared by King (J.C.S. 1930, 2374) by treating sarcosine in alkaline solution with methylisothiccarbamide hydriodide. Bergmann and Zervas have obtained diacetylcreatine from sarcosine and triacetylanhydroarginine (Z. physiol. Chem. 1928, 173, 80).

Creatine forms colourless transparent monoclinic prisms, containing 1 H2O, which it loses at 100°; it dissolves in 74.4 parts of water at 18°; in 9.810 parts of cold absolute alcohol, and is insoluble in ether (von Liebig, Annalen, 1847, 62, 257). Heat of combustion at constant pressure is 4,240 g.-cal. (Emery and Benedict, Amer. J. Physiol. 1911, 28, 307). The presence of urea, creatinine, or certain salts raises the solubility of creatine in water and alcohol (Nenbauer, l.c.). The aqueous solution is neutral, and has a bitter, harsh taste.

Creatine is decomposed by boiling baryta water into sarcosine, urea, and methylhydantoin

$$\begin{array}{l} \text{NH:C(NH}_z) \\ \text{NMe·CH}_2 \cdot \text{CO}_2 \\ \text{H+H}_2 \\ \text{CO(NH}_2)_2 \\ \text{+NHMe·CH}_2 \cdot \text{CO}_2 \\ \text{H} \end{array}$$

$$= CO(NH_2)_2 + NHMe \cdot CH_2 \cdot CO_2H$$

$$= CO(NH_2)_2 + NHMe \cdot CH_2 \cdot CO_2H$$

$$- CO(NH_2)_2 + NHMe \cdot CH_2 \cdot CO_2H$$

$$- CO(NH_2)_2 + NHMe \cdot CH_2 \cdot CO_2H$$

$$- NHe \cdot CH_2$$

$$+ NHMe \cdot CH_2 \cdot CO_2H$$

$$+ NHMe \cdot$$

(Neubauer, Annalen, 1866, 137, 294); or by mercuric oxide and water into oxalic acid and metbylguanidine; or by heating with soda-lime into ammonia and methylamine. Treated with mercuric acetate in presence of sodium carbonate it yields a-methylguanidinoglyoxylic acid,

NH2·C(NH)NMe·CO·CO2H,2H2O;

glistening flakes, m.p. 203-204° (Bauman and Ingvaldsen, J. Biol. Chem. 1918, 35, 277). When potassium hydroxide is added to an aqueous solution of creatine and silver nitrate until the white precipitate first formed just redissolves, the liquid solidifies to a transparent gelatinous mass immediately reduced on heating, or in the course of a few hours at the ordinary temperature. The addition of potassium hydroxide to a solution of creatine and mercuric chloride In order to extract creatine, the flesh is heated with its own weight of water at 60°, and the juice expressed. The flesh extract is then heated to coagulate the protein, and filtered; the fil- converted into creatinine by heating with dilute mineral acids for some bours (Dessaignes, | amino acid in the presence of ammonia, eq Jahresber. Chem. 1857, 544) or at 117° for 15 minutes (Beoedict and Myers, Amer. J. Physiol. 1907, 16, 362); or by boiling to dryness a solution of creatine to which has been added an equal volume of hydrochloric acid (Benedict, J. Biol. Chem. 1914, 16, 192); or by beating with water only in sealed tubes at 100° for 2-3 daya (Neubauer, Z. anal. Chem. 1863, 2, 33). The equilibrium between creatine and creatinine in aqueons solutions has been studied by Edgar and Shiver (J. Amer. Chem. Soc. 1925, 47, 1179). The ratio of the molecular concentration of creatinine to that of creatine in water at different temperatures and in various buffer solutions is given; thus in water the ratio is 0 54 at 25°; 2 89 at 100°; 36 72 at pn 1 (50°), and 1-03 at pa 6 (50°).

Smorodincev (J. Rnss Phys. Chem. Ges. 1915, 47, 1275) obtained a double salt of silver nitrate and creatinine, C. H.ON, AgNO, (decomposing 188-191°) by treating pure creatine with silver

mitrate in a slightly acid solution

Creatine forms soluble normal salts with the mineral acids (Dessaignes, Annalen, 1654, 92, 409), and readily soluble compounds with zinc chloride, C,H,O,N,,ZnCl₂, and cadmum chloride, C,H,O,N,,CdCl₂,2H,O (Naubauer, ibid. 1886, 137, 300). Creatine parents forms woully needles melting at 218-220° (decomp.) The mp is not depressed by admixture with creatinine picrate (King, J.C.S. 1930, 2374) The flatianais (v p. 417d) melts at 231° (Langley and Albrecht, J. Biol. Chem. 1935, 108, 729) When creatine in alkaline solution is treated with sodium nitroprusside and a persulphate a red colour is obtained; Pittarelli claims that this reaction is sensitive to 1 in 50,000 (A. 1928,

1121). Preating is not usually estimated directly; it is converted into creatinine (q.v.), and estimated as such; for the estimation of creatine in urine, v. Benedict (J. Biol. Chem. 1914, 18, 192; 1929, 82, 1; Folin, ibid. 1914, 17, 469). Methods for estimating creatine directly have been auggested by Walpole (J. Physiol. 1911, 42, 301), who makes use of the fact that a pale red colour is given by alkaline solutions of creatine and not by creatinine when a trace of discetyl is added, and by Lang (Z. physiol, Chem. 1932, 208, 273) who uses the violet colour formed from creatine and acctylbenzoyl in alkaline solution. Several methods have been adapted for blood filtrates, but it is still uncertain whether the chromogenic aubstance in blood is really creatine, see Peters and van Slyke (" Quantita-tive Chinical Chemistry," Baillière, Tindall & Cox. 1932).

a. Guanidinopropionic acid, alacreatine,

NH. C(NH)NH CHMe CO.H.

from alanine and methylisothiocarbamide. picrate forms needles, m.p. 187º (decomp.) (King, J.C.S. 1930, 2374).

a-aminocaprocyamine,

NH .: C(NH)NH CH(CO, H)CH, CHMe.,

from leueme and cyanamide (Duvillier, Compt rend. 1886, 103, 211; 1887, 104, 1290). Accord. ing to Duviller (l.c.) methyl or ethyl amino.

acids yield with cyanamide the corresponding creatinine. The creatine is formed only in the case of methylglycine and of \$\beta\$-methyl aminopropionic acid, the latter yielding methyl alacreatine, NH₂ C(NH)·NMe-CHMe CO₄H (Lindenberg, J pr. Chem. 1875, [ii], 12, 253). This generalisation is not confirmed by Gansser (Z. physiol Chem. 1909, 61, 16), who obtained the creatmine and not the creatine from a methylaminopropionic acid and sysnamide. methylaminopropionic acca and cynnamic, but prepared β methylguanidinopropionic accd, NH₂C(NH)NMe[CH₂]₂CO₂H,H₂O, mp p 201–202°, from cynnamide and β methylamino propionic acid, and γ methylamino propionic acid, and γ methylaminofoliolity acid, NH₂C(NH)NMe[CH₂]₂CO₂H, mp. 307°, from eyanamide and y-methylaminobutyne

Formaldehyde and creatine yield the compound CaH, O, N, 2H, O, decomposing at 250°, and forming a dibenzoyl derivative,

C.H.O.N.Bz.

m p. 265-268° (Jaffé, Ber. 1902, 35, 2896). Phthalyldicreatine.

C.H.[CO NH·C(NH)NMe·CH,·CO.H], hae m.p. 212° (Urano, Bait. Chem. Physiol. Path, 1907, 9, 183). Diacetylcreatine,

NH.C(NHAc)NMe CH, CO,Ac, has m p. 165° (Erlenmeyer, Annalen, 1895, 284,

M. A. W. and W. V. T. CREATININE. Methylglycocyamidine, -NMe CH.

C:NH NH CO

discovered by von Liehig in human unne (Annalen, 1847, 82, 268), is also a constant constituent of the urine of horses, cows, dogs, pigs and rabbits; the amount of creatmine in the tissuea is uncertain as its determination is attended with difficulties owing to the ease with which creatine is converted into creatinine by post morten autolysis. This accounts for the conflicting reports of different investigators on the creatinine content of muscle which probably contains only a very small amount if any, According to Folin and Denis (J. Biol. Chem. 1914, 17, 487) norms buman blood contains 1 mg, creatinine and 6-8 mg. of creatinine and creatine per 100 g blood: but see below under Estimation of Creatinine. In domestic animals it is about the an isomer of creatine, forms aparingly soluble same; in hirds 0-1 mg, creatinane but 11 mg prisms. 1t is prepared from alaume and creatinine and creatina. 1t is probably a normal cyanamide (Baumann, Annalen, 1873, 187, 83) or constituent of all soils (Shorey, J. Amer. Chem. The Soc. 1912, 34, 99; Sullivan, ibid. 1911, 33 King. 2035) and vegetable matter (Oshima and Am zumi, J. Coll. Agric. Tokyo, 1914, 6, 17)-

The homologues of creatine are prepared by Under normal conditions, the exception of the action of cyanamide on the corresponding creatinuse is constant from day to day. The

amount excreted by an individual per day varies from 7.5 to 10 mg. creatinine nitrogen per kg. body weight and is roughly proportional to his muscular development; it is not affected by diet or exercise. Creatinine is, therefore, generally regarded as an end product of endogenous metabolism, most probably of creatine catabolism (Shaffer, Amer. J. Physiol. 1908, 23, 1; Folin, *ibid.* 1905, 13, 45, 66, 117; Pekelharing, van Hoogenhuyze and Verploegh, Proc. K. Akad. Wetensch, Amsterdam, 1905, 8, 363; Klercker, Beit. Chem. Physiol. Path. 1906, 8, 59; Folin and Denis, J. Biol. Chem. 1914, 17, 500; Harding and Gaebler, *ibid.* 1922, 54, 579; Hodgson and Lewis, Amer. J. Physiol, 1928, 87, 288). Creatinine administered to animals is rapidly excreted almost quantitatively (Foster and Fisher, J. Biol. Chem. 1911, 9, 359; Towles and Voegtlin, ibid. 10, 479). Under ordinary circumstances no creatine is excreted by adults. Some normal women, however, all infants and young children, and adults in starvation and certain pathological conditions excrete some creatine (Denis and Minot, ibid. 1917, 31, 561; Folin and Denis, ibid. 1912, 11, 253; Rose, ibid. 1912, 10, 265; 1917, 32, 1; Gamble and Goldschmidt, ibid. 1919, 40, 199).

Creatinine is extracted from urine by evaporating to one-third the original volume, decanting from the salts that crystallise out, precipitating with lead acetate and filtering, removing the lead from the filtrate and precipitating the creatinine with mercuric chloride as the sparingly soluble double salt. This is separated, decomposed by sulphuretted hydrogen, and the creatinine crystallised from the filtrate as the hydrochloride (Maly, Annalen, 1871, 159, 279; or by precipitating it as creatinine potassium picrate with subsequent decomposition of the double salt (Folin, J. Biol. Chem. 1914, 17, 463; Benedict, ibid. 1914, 18, 184)

For extraction from the body tissues and fluids, see Costantino (Chem. Zentr. 1915, 11, 287, from Arch. farm. sper. 1915, 19, 254).

Creatinine bears the same relation to creatine as hydantoin to hydantoic acid

and is prepared by boiling creatine with dilute mineral acids for some hours (Liebig, Annalen, 1847, 62, 628) or by heating at 117° for 15 minutes under pressure (Benedict and Myers, Amer. J. Physiol. 1907, 18, 362, see, however, J. Biol. Chem. 1914, 17, 363; Folin, ibid. 469), or by boiling down to dryness with an equal volume of N-HCl (Benedict, ibid. 1914, 18, 192), or by heating creatine with anhydrous zinc chloride (Edgar and Hinegardner, ibid. 1923, 56, 1).

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Creatinine can be prepared synthetically by heating guanidine carbonate with sarcosine at 140-160° (Horbaczewski, Maly's Jahresber. Tierchem. 1885, 15, 86). A quantitative yield is obtained by heating N-methylglycocyamine with an organic acid (G.P. 281051, from J.S.C.I. 1915, 34, 637).

By heating creatinine with silver nitrate and baryta in excess on the water-bath for about an hour a nearly quantitative yield of methylguanidine can be obtained (Ewins, Biochem. J. 1916, 10, 104). By oxidation with mercuric acetate it yields a-methylguanidinoglyoxylic acid (Bauman and Ingvaldsen, J. Biol. Chem.

1918, 35, 277).

Creatinine forms anhydrous monoclinic prisms soluble in 11.5 parts of water or 625 parts of absolute alcohol at 17° (Joppelius and Pommerehne, Areh. Pharm. 1896, 234, 380), and separates on slow evaporation of dilute solutions in square plates containing 2H₂O (Salkowski, Z. physiol. Chem. 1880, 4, 133). It is a weak base, displacing ammonia from its salts, and forming soluble salts with the mineral acids (Liebig, l.c.), the picrate C₄H₇ON₃·C₆H₃O₇N₃, m.p. 212-213°, forms sparingly soluble yellow needles; the acid picrate C₄H₇ON₃·2C₆H₃O₇N₃, has m.p. 161-166° (Mayerhofer, Wien. Klin. Woch. 1909, 22, 90); the flavianate, solubility 0·198, melts at 250° (decomp.) (Kossel and Gross, Z. physiol. Chem. 1924, 135, 168); the tartrate (C₄H₇ON₃)₂C₄H₆O₆ decomposes at 207-209°; the oxalate (C₄H₇ON₃)₂·C₂H₂O₄ is sparingly soluble in alcohol (Poulsson, Chem. Zentr. 1904, ii, 30). Creatinine forms characteristic double salts, with platinic chloride,

(C4H7ON3,HCI)2·PtCl4,

orange-red plates, m.p. 220-225°; with auric chloride, C₄H₇ON₃·HCl·AuCl₃, yellow crystals, m.p. 170-174°; with potassium picrate, C₄H₇ON₃·C₆H₃O₇N₃·KC₆H₂O₇N₃, lemon-yellow prisms, 100 parts of water dissolve 0·1806 part of the salt at 19-20°; with zine chloride, (C₄H₇ON₃)₂·ZnCl₂, almost insoluble in alcohol (Heintz, Jahresber. Chem. 1847, 48, 883); with mercuric chloride,

4(C₄H₅HgON₃·HCl)·3HgCl₂,2H₂O

(Johnson, Proc. Roy. Soc. 1886, 43, 493).

Creatinine is converted into creatine by the aetion of cold dilute ammonia or calcium hydroxide solutions; or on treatment with iodine (Reichardt, Pharm. Ztg. 1911, 56, 922); boiling baryta solution converts it into ammonia and methyl hydantoin; with alkaline oxidising agents it yields methylguanidine and oxalic acid, but is comparatively stable towards acid permanganate solutions (Jolles, Ber. 1902, 35, 160).

The presence of creatinine in dilute solution or in urine can be detected by (1) Weyl's test (Ber. 1878, 11, 2175), which consists in the production of a ruby-red coloration, changing to yellow when, to a solution of creatinine, a few drops of 10% sodium nitro-prusside and the same quantity of 10% sodium hydroxide or carbonate solution are added. On acidifying the

1 "Flavianicacid" is the name applied by the authors to naphthol yellow S, 1-naphthol-2: 4-dinitro-7-suiphonic acid.

solution with acetic acid the colour is discharged,] turning to green or hlue on boiling due to the been employed for the estimation of creating formation of Prussian blue (Salkowski; Colssanti, Gazzetta, 1887, 17, 129). Guareschi with alcoholo 3:5 dinitrobenzoic acid and 6% (Ber. 1888, 21, Ref. 372) has shown that this sodium hydroxide. Creatine, glucose and ketoms reaction is given by compounds other than creatinine containing the —CO CH — group.

(2) Jates test (2, physiol. Chem. 1886, 10, that the chromogenic substance is not creating. 399) is the production of a deep red colour, when pieric acid and an excess of sodium hydroxide solution are added to the liquid containing the creatinine. It is now generally accepted that the red colour in Jaffé's reaction is not due, as suggested by Chapman (Chem. News, 1909, 100, 175), to sodium salts of picramic acid and diaminonitrophenol formed hy reduction of the pieric acid by the creatinine, as the red compound has been isolated and shown to be a red tautomer of creatmine picrate although its structure is still under discussion (Greenweld, J. Biol. Chem 1924, 59, 601; 1928, 80, 103; 1930, 86, 333; Anslow and King, J.C.S 1929, 1210)

Estimation of Creatining.-The older methode for the estimation of creatinine were based upon the isolation of the hase in the form of its sparingly soluble double salt with mercure or zinc chloride, and either weighing the precipitate zinc carorae, and other weight the preophato or estimating the metal or the nitrogen in the eompound (Neubauer, Annalen, 1861, 119, 35; Sakkowski, Z physiol, Chem. 1886, 10, 113; Grocco, Chem Zentr. 1887, 17, Kolisch, ibid. 1895, 1, 814, Ladd and Bottenfield, Amer. 1895, 1, 814, Ladd and Bottenfield, Amer. Chem. J 1898, 20 869). Edlefsen (Chem. Zentr. 1909, 1, 108) recommends precipitating the sulphate in the precence of alcohol and ether, duscolving it in water, and titrating the solution with standard barium hydroxide solution, using phenolphthalein as indicator.

The method generally adopted for estimating creatinine in urine is Folin's colorimetric method. based on Jaffé'e reaction (Z. physiol, Chem, 1904, 41, 223; J. Biol. Chem, 1914, 17, 469), If creating is also present in the using the total creatizate is determined after converting the ereatine into creatinine hy heating with acid, Subtraction of the preformed creatinine, determined separately, gives the creatine as creatin ine. The conversion of creatine into creatinine may be carried out by (1) boiling with a saturated solution of pieric acid, and this method must be used if glucose is present (Folin, Ic.); (2) autoclaving with saturated pieric acid (Folin, Ie.); or (3) holling down to dryness with hydrochloric acid in the presence of metallic lead (Benedict, J. Biol. Chem. 1914, 18, 191). It is important that pure picuc and should be used for the colorimetric determination of creatinine by any method involving Jaffe's reaction (Folin and Doisy, ibid. 1917, 28, 349). Benedict gives outable methods for the purification of pieric acid for this purpose (ibid. 1922, 54, 239; 1929, 82, 1).

The presence of creatining in blood stall remains to be established with certainty. Whilst there is a chromogenic substance which can be estimated by Folm's method (Folin and Wu, told. 1919, 38, 81) it is probably in the gives an aurichloride, m.p. 170-171, and main not creatinine (Behre and Benedict, dimethylcreatinine gives an aurichloride, mp ibid. 1922, 52, 11; and see below).

More recently a new colormetric method has in urme This consists in treating the unit with alcoholic 3:5 dinitrobenzoic acid and 6% that the chromogenic substance is not creatinine (Benedict and Behre, ibid. 1936, 114, 513; Langley and Evans, ibid. 115, 333).

An isomeride of creatinine, alacreatining

was prepared (Ber. 1873, 6, 1371), and other isomerides have been described by Korndörfer, Arch. Pharm. 1904, 242, 620; Schenck, bid. 1910, 248, 376; 1911, 249, 463; Schmidt, bid. 1910, 248, 585. Johnson Micolet (J. Amer. Chem. Soc. 1915, 37, 2417) ohtained two methylglyeocyamidines isomeno with creatinine by the action of methylamine on the ethyl ester of benzoylpseudoethylthichydentoic acid and debenzoylating.

Two isomerio nitrosocreatinines were obtained by Dessaignes (Annalen, 1856, 97, 341) and Märcker (1814 1865, 133, 305), by passing the nitrous finmes from the action of nitric acid on arsentous oxide into an acid solution of creatinine. The less soluble a nitrosocreatinine, C. H.O.N. decomposes at 210°, forming a colourless houd that quickly solidifies to a brown mass from which a new base, C,H12OaN10 can be er-tracted; it forms a crystalline nitrate, hydro chlorde and platnichloride, and yields the bromo-derivative C,H,O,N,Br. The more soluble B,nitroscerationne, C,H,O,N, melts at 195° to a brown liquid and decomposes at 220°.

According to Kramm (Chem. Zentr. 1898, L 37), the yellow colour formed by the interaction of sodium nitroprusside and sodium hydroxide on creatmine (cf. Weyl, Ber. 1878, 11, 2175), is due to the formation of a nitrosocreatining, C4HaO2N4, which can be isolated as a colourless crystalline precipitate by adding acetic acid to the yellow alkaline solution and absking vigorously. According to Schmidt and Bennig (Arch. Pharm. 1912, 250, 345) this is an oxime; they find that on treatment with sodium nitnie in nitric acid colution creatining yields a mixture of methyl hydantom oxime, m p. 193-194°, and creatmane oxime, which begins to discolour at 250° hut does not melt.

The following acyl and alkyl derivatives of creatinine have been prepared : Benzoylcreatinine, C₁₁H₁₁O₂N₃, pale yellow needles, mp. 187° (Urano, Beitr, Chem. Physiol. Path. 1907, 9, 183); methyl, dimethyl, and trimethyl creatinine and salts (Korndorfer, Arch. Pharm. 1904, 242, 641; Kunze, Arch. Phsrm. 1910, 248, 578) Methylcreatinine

128-129°, and a platmichloride, m p. 177-179'.

creatinine is

it reacts as a tertiary base; ethylcreatinine

(Neubauer, Annalen, 1861, 119, 50), see Henzerling (Arch. Pharm. 1910, 248, 594; the platinichloride crystallises in monoclinic plates, m.p. 197-211°; ethylcreatinine hydriodide, m.p. 217-219°; diethylcreatinine platinichloride 201 -202°).

For various other substituted derivatives of creatinine, see Duvillier (Compt. rend. 1882, 95, 456; 1883, 96, 1583; 97, 1486; 1885, 100, 916; 1886, 103, 211; 1887, 104, 1290) and Greenwald (J. Amer. Chem. Soc. 1925, 47, 1443). M. A. W. and W. V. T.

CREOLINE. A coal tar disinfectant prepared from coal tar creosote, contains 45-60% neutral oils, 10-15% phenols, 30-40% rosin

CREOSOL (4-hydroxy-3-methoxy-1-methyl-

benzene) v. CREOSOTE.

CREOSOTAL (creosote carbonate) is a mixture of carbonates of the various phenols contained in creosote, chiefly creosol, guaiacol, and cresols.

It may be prepared by heating creosote with a benzene solution of phosgene under pressure, but is generally produced by passing a stream of phosgene into a solution of creosoto in aqueous caustic soda. Creosotal separates as an oil and is washed with dilute caustic soda solution and then with water. It contains the equivalent of about 90% of creosote.

Creosotal is a clear, colourless or yellowish, viscid liquid, insoluble in water but soluble in alcohol, ether, chloroform, benzene, (Aufrecht, Pharm. Ztg. 1908, 53, 480). benzene, etc. It is neutral, has little odour and only a slight taste of creosote. Crystals of guaiacol carbonate may settle from samples after long standing or on freezing. Boiling with alcoholic caustic alkali gives creosote and alkali carbonate.

It has to some extent replaced creosote in treatment of pulmonary tuberculosis,

bronchitis and pneumonia. CREOSOTE. Tars obtained by the destructive distillation of wood and of coal in retorts, coke ovens, or blast furnaces, when subjected to distillation yield fractions techni-cally known as "creosote," and which find extensive and important uses in the industrial arts, the most important type being that derived

from gas works or coke oven coal tar.

The term "kreosote" was originally applied to the product obtained from wood tar. The increase in production and distillation of coal tar with the extension of the use of the creosote oil so obtained for the preservation of wood resulted in the terms creosote and creosote oil being used commercially as meaning the heavier

According to Kunze the constitution of dimethyl- I prefix to indicate their origin, as "blast-furnace creosote." In pharmacy the term creosote is still retained for wood creosote.

WOOD-TAR CREOSOTE.—The tar derived from the distillation of beechwood and other hard woods, when subjected to further distillation, yields certain fractions that are heavier than water. These are agitated with an aqueous solution of caustic soda, which is then separated from insoluble oils, boiled with free access of air, in order to oxidise various impurities present and to distil off hydrocarbons, and then decomposed with dilute sulphuric acid or carbon dioxide. The crude creosote which separates is again submitted to treatment with alkali and acid, and finally distilled, the fraction distilling between 200° and 230° being collected separately.

Wood-tar creosote, when freshly prepared, is a colourless, transparent liquid of an oily consistency, and which retains its fluidity at a very low temperature; its sp.gr. varies from 1.037 to 1.087; it distils between 200-230°, and dissolves in about 200 parts of water; its odour is strong and penetrating, resembling that of wood smoke; it has a high refractive index (1.51) and burns with a luminous smoky flame. It is neutral or only faintly acid to litmus. It is powerfully antiseptic, but does not coagulate a bumin; is not so caustic as carbolic acid and relatively less poisonous. Wood creosote is essentially a mixture of the ethers of the polyhydrio phenols, chiefly guaiacol (o-methoxy phenol, b.p. 205°) and creosol (2-methoxy-4-methyl phenol, b.p. 221°); whilst phenol, o-, mand p-cresols, o-ethyl-phenol and xylenols may be present in varying amounts. The presence of the dimethyl ethers of pyrogallol (b.p. 262°) and of methyl-and propyl-pyrogallol (b.p. 265° and 285°) has been identified in samples. Dimethylguaiacol (b.p. 230°) and propylguaiacol (b.p. 241°) are present in insignificant amounts. The latter is objectionable in the medicinal grade, since a single drop on the tongue causes bleeding; its presence may be recognised by the blue coloration produced with baryta water.

The percentage composition of beechwood and oak creosotes, freed from hydrocarbons, is given by Béhal and Choay (Compt. rend. 1894, 119, 166) as follows:

ı		Beech.	Beech.	Oak.
	Distillation temp.			
ĺ	°C	200-2209	200-210	200-210°
Į	Specific gravity .	1.085	1.085	1.068
	Mono-phenols	39-0	39.0	55.0
ı	Guaiacol	19.7	26.5	14.0
1	Creosols and homo-			
i	logues	40.0	$32 \cdot 1$	31.0
ı	Loss	1.3	$2 \cdot 4$	_
	Guaiacol Creosols and homo-	19·7 40·0	26·5 32·1	14.0

Wood creosote is miscible in all proportions with alcohol, ether, glacial acetic acid, benzene, carbon disulphide and light petroleum spirit. It dissolves in concentrated sulphuric acid to a red liquid, which slowly changes to purpleviolet; it is violently attacked by nitric acid, is soluble in aqueous solutions of alkali hydroxides. and forms a crystalline compound with potassium distillates from gas works and coke oven tar, hydroxide, but not with sodium hydroxide. On the oils from other tars being distinguished by a the other hand, it is practically insoluble in

sp gr. at 15 5° not helow 1.070, commencing to distil at about 200°, and yielding not less than 95% between 200° and 230°; 2 c c. should require 10-18 c.c. N NaOH to produce a clear solution, and the liquid should remain clear on dilution with 50 cc. of water (limit of hydrocarbons, and of hases). Owing to the demand for guaracol for pharmaceutical purposes, much of the wood creosote now sold is frequently found to have been deprived of part or the whole of the guaracol it contained. Methods for the determination of guaracol in wood creosote are given by Béhal and Choay (Compt. rend. 1893, 116, 197); Kebler (Amer. J. Pharm. 1899, 71, 409). Woodtar creosote is distinguished from coal-tar phenois by its reaction with an ether-alcohol solution of nitrocellulose (equal parts), no

coagulation heing produced. Wood creosote is used as an antiseptic and analgesio material (external), and internally in the treatment of tuberculosis by preparations such as creosote carbonate and the lactic acid and valeriance esters. Also employed in the preservation of mine timbers, in lignin removal from jute fibre, and in oro flotation. Bubo (BP 257151) employs aq ethyl alcohol at 50°C for extraction of phenols from wood tar The alcoholic extract of brown coal tar has been placed on the German market under the name Frest—this product has apgr. 1-01-1-04, visconity at 20°C, 5-10° Engler, flash-point 100-110°, constituents sol. m 38° Be NaOH, 60-90%, colour black. Gracio (G P. 232657) extracts with alcohol in a continuous column

apparatus—against saving in chemicals over older methods is to be placed a loss of only 0.3-0.5 parts alcohol on 100 parts of tar. Coat. Tar Creosore.—The creosote oils derived from coal tars (cf. Coat Tar) may be divided into two main classes-light creceote heing the fraction distilling mainly between 190° and 235°, and ordinary creosoto oil, chiefly employed for timber preservation, and distilling mainly between 200° and 300° to 350°C. The light creosote normally consists of the middle or agai creacte normally consists of the module or aerholic oil fraction obtained in the primary destillation of the tar, the crystallisable usphths-lene having been removed by cooling the oil to atmospherio temperatures. If the oil has been treated for the recovery of plenols, the residual content of these compounds may be 3-10%, If the oil is to be employed for disunfectant and thio indigo series, has been recovered from the tar acids, water 0.7%, and carbon 85.75, hydro-265-275° fraction, and indole from the 240-gen 7 92, oxygen+nitrogen 5-14, sulphar 0 49%.
260° fraction (G.P. 223304). Formerly the The gross calorific value of horizontal retort tar oil distilling between 300° and the end of the oil has been given as 9 900-50T cals, where T distillation was separately collected, cooled, is the percentage of phenois present, and the clear oil (filtered anthracene oil or green efficient of expansion is 0 00075 per deg. C.

strong ammonia. Official in Brit. Pharm. 1932, [oil] separated from the crystalline crude anthra cene. The crude anthracene was purified to yield a high-grade anthracene and carbarole. and the fluid oil was sold for the preservation of timber by hrush treatment (carbolineum infra). A fall in the demand for anthracene, owing to the anthracene derivatives for the dyestuffs industry heing more simply obtained synthetically, coincided with a growing difficulty in obtaining anthracene of low paraffin in nonadecane) content from many modern types of tars, and at present in the United Kingdom the authracens oil fraction is frequently run

direct to the creosote storage wells. Composition: So many varying factors in fluence the composition of creesote oils that it is impossible to give more than the following general list of some of the more important compounds that have been found to be present in the samples examined : hydrocarbons-naphths. lene, a and \$ methylnaphthalenes, 16. 2.3. 2.6-, 27, dimethylnaphthalenes, anthracene and & methylanthracene, phenanthrene, dipheny and its methyl and dimethyl homologues, ace naphthene, 45-henzoindane, fluorene together with varying amounts of saturated and un saturated hydrocarbons of the aliphatic sens. oxygen containing compounds-phenol homologues, chiefly crosols and xylenols, a and \$ naphthol, diphenylene oxide and its methyl nitrogen containing compoundsdenvative: pyridine and its methyl and dimethyl home logues, anime and toluidine, quinoline and isoquinoline and their methyl and dimethyl homologues, a. and β-naphthylamine, acridine, carbazole, indole and its methyl homologues; sulphur-containing compounds—phenylthiol, thionaphthene, thiodiphenylene. The oils from vertical retort tars contain a higher proportion of aliphatic hydrocarbons than those obtained from horizontal retort and coke over tars, and this is indicated by the lower specific gravity of the close-cut fractions obtained by redistillation.

Physical Properties: The latent bests of evaporation of tar distillates have been deterowaporation of ter distillates have been defermined by Weiss (Ind. Eng. Chem. 1922, 14, 72) fraction 249-299°C., 146 B.Th.U. per bl., 229-543°C, 137. 35-32°C., 132; 332-435°C., 117. Speam heat of tradistillate of 34±00 at 16-90°C. (International Critical Tables), 04 at 40°O at 80°O at 180°C. (Revebs. Chem. u. Vyasarrichh. 134; 70, 280°C.) If the oil is to be employed for disinfectant and [Krebs, Gas- u. Wasserfach, 1930, 73, 821, sheep dpm annufacture the creed content me, the color of the properties to 40% (as-called crevylor crecostes). The 1-049 at 20°C, 16,544 h.Th.U./lb, gress, 15,530 morthage recosto contains the oils dathling from B.Th.U./lb, net; vertical retort tar-oil, apgrate to the tarbetween the end of the ear-lose oil or 1-016 at 20°C, 16,540 h.Th.U./lb, gress, 15,730 maphthalene oil fractions and the pitch, and B.Th.U./lb, net; blast furnace tar-oil, apgrate and the properties of the color of the properties of the p

finds its largest application in the preservative treatment of timber, especially railway sleepers, telegraph poles, and marine piling, against fungi, white ants, and marine borers respectively. The resistance to mechanical forces is also improved by creosoting. The largest consumer, the United States of America, uses annually upwards of 100 million Imperial gallons, 180 million gallons being used in the peak year, 1929. In 1935 the U.S.A. production was 90 million gallons; in 1936 34 million gallons were imported, of which 20 million gallons came from the United Kingdom. Formerly the U.S.A. absorbed 50% of the British production, but from 1929 the British export figure to U.S.A. fell owing to the increased American production, and reached a minimum at 9 million gallons in 1932. No official figures for the United Kingdom are available, but the production of creosote oils of all grades for the year 1935 and 1936 is estimated at 80 and 85 million gallons respectively, the use figure for timber treatment at 12-14 million gallons per annum, and the total export figures for 1935 and 1936 are 23½ and 36½ million gallons respectively.

For estate timber, fluid creosotes may be applied by brush or spray, or the posts, etc., may be immersed in the oil at 180°F, contained in open tanks, and allowed to cool while still immersed. The more effective method is to apply the creosote oil under pressure, steel cylinders 75 to 105 ft. in length by about 7 ft. in diameter being employed. In the Bethell "full-cell" process, the timber is subjected to a vacuum of 22 inches for 30 minutes, and the oil at a temperature of 180-210°F. is then introduced without breaking the vacuum until the cylinder is full. The pressure is then raised to 100 to 200 lb. per sq. in., and when absorption is complete the surplus oil is run off and a vacuum may be applied for 15 minutes (Lowrie modification). Green timber may be conditioned by a preliminary heating for 24 hours in the oil under vacuum, maximum temperature 200°F., to remove the surplus water (Boulton process). In the Rueping "empty-cell" process the timber is subjected to an initial pressure of 10 to 60 lb. pcr sq. in., and the oil is then introduced into the cylinder, the pressure being maintained constant. The pressure is then raised to 80 to 120 lb. per sq. in. to give maximum injection. The amount held in the timber is then reduced to the required figure by expansion of the air initially introduced and by the application of a high vacuum for a short period. Absorpt on figures are of the order of 15 to 20 lb. per cu. ft. and retention 7 to 10 lb. per cu. ft. In be Card process, zinc chloride and creosote re employed, typical absorption figures being 2 lb. zinc chloride and 3 lb. of creosote per cu. ft.

The quantity of oil absorbed depends upon the species, growth and size of the timber, the creosoting process employed, and the admissible cost. Penetration of the oil may be assisted by incising, that is, by making a number of small cuts over the surface of the timber parallel to the grain and about $\frac{3}{4}$ in. deep by 1 in. long. The usual figures lie between 6 and 20 lb. per cu. ft.

COMMERCIAL USES AND GRADES.—Creosote to 25 years and upwards, dependent upon the desits largest application in the preservative conditions of employment, such as climate, estment of timber, especially railway sleepers, location, mechanical stresses, etc.

The desirable properties in a creosote oil are toxicity, permanency and power of penctration and waterproofing. Considerable laboratory work on the relative toxicity of the constituents and fractions of coal tar creosote has not led to conclusive results. The more important standard specifications are as follows: B.S.. 144/1936, Type A, calls for an oil sp.gr. ³⁸/₂₀ 1·010–1·065; completely liquid on warming to 38°C., and on cooling remaining completely fluid after standing 2 hours at 32°C.; water content not more than 3%; yielding on distillation to 205°C. maximum 6%, to 230°C. maximum 40%, to 315°C. maximum 78 w/w%, residue above 315° soft and not sticky; phenols 5–16%; matter insoluble in benzole maximum 0.4%. Type A2 admits an oil of sp.gr. 38 0.995–1.065, the other clause being as before, except that no maximum is specified for the phenol content and the maximum allowable distillate at 315° is 85%. Type B covers creosote from coal-tar made in Scotland, the details being as for Type A2. Creosote from low temperature tar is admitted under type A2, with a minimum specific gravity 0.935, and blast furnace creosote under type B with similar limits. Creosote required for brush application should be fluid on cooling and standing at 15.5°C. for 2 hours. For the distillation test a standard side arm flask and thermometer graduated for 100 mm. immersion are employed. The American Railroad Engineers Specification for No. 1 grade oil is sp.gr.38 not less than 1.03; water maximum 3%; matter insoluble in benzole maximum 0.5%; distillate on a water-free basis up to 210° maximum 5%, to 235° maximum 25%; coke residue maximum 2%. The distillation test uncorrected for barometric pressure is performed in a special standard flask, a total immersion thermometer being employed. The American Railroad Engineers Association oil in general meets the continental specifications, except that in Scandinavia and Germany the maximum distillate to 235° is limited to 20%, and a phenol clause is included (Scandinavia, 3-8%).

Commercial oils are sometimes diluted with oil-gas tar and its distillates, or with mineral oils; these mixtures have a lower conservation value. In America coal-tar creosote solutions are employed, of which at least 80% is a distillate from coal-gas or coke-oven tar, and the remainder refined or filtered coke-oven tar. The standard American Wood-Preservers Association specification is sp.gr. 18-105-1-12; water maximum 3%; distillate on a water-free basis to 210° maximum 5%, to 235° maximum 25%; matter insoluble in benzole maximum 2%; coke residuc maximum 6%. Literature.—Proceedings of British and of American Wood-Preservers Associations.

Carbolineum is a fluid, high-boiling, tar-oil, distilling maximum 10% at 250°C; sp.gr. at 20° 1.08-1.11; phenols maximum 10%; water maximum 1%.

usual figures lie between 6 and 20 lb. per cu. ft. The A.R.E.A. Specification for antbracene oil The life of the timber is raised from 3 to 6 years covers an oil of sp.gr. 185 1-09-1-13; distilling

CREOSOTE.

not more than 1.5% to 235°, 16.5% to 300°, | tion still amounted to 16,000 tons. Crecacte oil residue maximum 2%.

Other outlets for creosote oil are:

Disinfectants and Sheep Dips.—An oil onitable for disinfectant manufacture should have op gr. 0 95-1 00 and distil mainly hetween 200° and 300°. The oil should he free from crystallisable naphthalene at 5°C., and the tar acid content (mainly cresols) will vary with the type of product required. Oils yielding pink emulsions are not favoured for sheep dips.

Fuel Oil for Furnace Work, particularly in the metallurgical and glass industries, is fed through atomising hurners operated by compressed air, and should normally have specific gravity not exceeding 1 07 at 38°C ; water content not ex-

ceeding 1 0%; flash point not helow 150°F. (cf. B S 503/1933). The above oil may require steam heated storage and pipe lines, which serve not only to prevent separation of naphthalene or anthracene, but also to lower the viscosity and improve atomisation in the humer Alternatively, the creosote may be mixed with oil gas tar, de hydrated coal-tars, or with petroleum fuel oils In hlending with oils of epecific gravity below unity, the specific gravity of the mixture requires careful attention to avoid mixtures of unit specific gravity. Owing to the high carbon and low bydrogen content, the heat lost in flue gasea is low. The gross calorific value is of the order of 16,600 B Th.U. per lb. The removal of phenols from tar-oils raises the calorific value by about 35 B.Th.U. per lb. for each 1% removed On the Continent a mixture of 80% tar oil and 20% pitch has been employed, specification heing fiash point (Pensky-library, minimum 75°C. (open); viscosity maximum 8º Engler at 20°; no deposit at 8°C. for 2 hours; free carbon maximum 4%; calorific value 8,500 g -cal minimum.

Spark Ignition Engine Fuel.-Considerable attention has been directed to this potential market. Creosote oil can be used with a greater thermal efficiency than petrol, but preheating of the oil entering the cylinder is necessary. and the employment of a volatile hydrocarbon fuel is necessary for starting up the engine, and for use when idling. Advance has been made in the design of suitable dual carburettors, but the results have not fulfilled expectations (Inst. Cas Eng. Report, 1934) The creosote oil should he free from crystallisable naphthalene at the prevailing atmospheric temperatures, should distil almost wholly below 265°C, the water content should not exceed 0 50%; phenols maximum 10%; and the coke value (Conradson test) should not exceed 0 1%. An alternative fuel is obtained by admixture of the creosoto oil with up to 20% of crude benzole or sulvent naphtha. These fuels have a high anti knock valua and are suitable for engines of high compression ratio, but crank case dilution is high and plug gumming troubles are difficult to

Compression Ignition Engine Fuel -Creosoto

45% to 355°; water maximum 0.5%; coke has a high spontaneous ignition temperature (440-490° in oxygen) and modifications in engine design are necessary to ensure satisfactory running. Improved nozzles and flams plates to give fine atomisation have led to success. The compression should not be less than 480 lb. per sq. in , the fuel admission must be advanced and pilot jet ignition employing a petroleum distillate normally 2 to 4% of the total fuel at full load is used. Although slight modification to slow. apeed Diesel engines may overcome difficulty of high ignition temperature, this property, together with long ignition delay of tar fuel oils, renders them unsuitable for employment in the modern high speed engine of the Dresel type. It has been chown that the phanols are partly responsible for the high ignition temperatures, but their removal does not reduce the ignition tempers ture to a figure comparable with that of the petroleum Diesel oils. Blending with shala oils, parathin oils, and synthetic oils (Glucksuf, 1936, 29, 697) has given promising results, and the cetene number of creosote oils has been raised from about 15 to a satisfactory figure of the order of 45 by the addition of 30% and upwards of these oils. Experimental work employing added primers, substances of low spontaneous ignition point such as trinitro resorcinol, methyl nitrate, hydrazine perchlorate, nitroxyl cthylene chlorhydrin, etc., has not yet yielded useful results, partly due to cost (cf B.P. 436027, 461320; C.P. 574678, 612073) Successful results are claimed by the employment of an insulated catslyst contact mass (thorsum oxide or platinum on asbestos wool or vanadium pentoxide supported on silica gel) hult into the pre-combustion chamber of the engine in the path of the atomised oil. It has been shown that tar-oils which give s bloore ignition temperature in oxygen of 449-470° give ignition temperatures of , 165-180° in the presence of platinum on asbestos, the normal figure for American gas oil being 248°C. (Cluckauf, 1932, 68, 980). Developments in design, including better atomission and increased temperature of fuel either within or prior to the combustion chamber, may produce an engine that is less fuel-sensitive. A suitable oil may have the following characteristics: fluid at prevailing atmospheric temperatures, water below 1%, and preferably 6.5% or lower; distillation test minimum 60% at 300°C; matter insoluble in xylol maximum 0.2%; cole residue (Conradoon test) maximum 30%; ach maximum 0 02%; flash point minimum 65°C; viscosity maximum 3° Engler at 20°C. Some users epecify maximum phenols content 10%. maximum sulphur content 0 5%. and colorino maximum 0 02%.

Benzole Wash Oil to he employed for the

recovery of benzole from town and coke oven gas should have opgr. at 15°C. 101-103; datatilation test ni at 200-21°C. not less than 70% at 300°C.; phenois normally 7-10% hut it is generally agreed the lower the better; water maximum 1%. The crystallisable con oils were used during the period 1014-1919 in this control and in Sermany for large stationary cooling to oil to 55°F, or may be hanted to Desel engines. In 1029 the German consump - 7% on cooling to 40°F, the fraction dutilling to

824) gives distillation test nil at 200°C., 90% helow 300°, the distillation curve to he as nearly linear as possible. The presence of anthracene is undesirable. Viscosity of fresh oil is 1.5° Engler at 20°C., the spent oils have a viscosity of 8° and upwards. For data on absorptive capacity of oils, see "Motor Benzole" (Hoffert and Claxton).

Hydrogenation.—By treatment at elevated temperatures (440-500°C.) and in presence of hydrogen under high pressure (200-265 atmos.) the rate of reaction heing accelerated by catalysts (molybdenum sulphide, or oxide, zinc oxide, and magnesium oxide on active carbon or silica gel), creosote from high and low temperature tars undergoes hydrogenation-cracking to yield lower boiling spirits of increased hydrogencarbon ratio. Crude low temperature tar may also be used as raw material. Tar-oils of appropriate distillation range (i.e. not distilling above 300-325°) may be treated directly in the vapour phase, hut tars need preliminary treatment in the liquid phase and are fed to the plant with the coal where this is heing treated. The Billingham plant, which started operations January, 1935, is stated to have heen designed to handle annually 10,000 tons low temperature tar, 40,000 tons coal tar creosote, 100,000 tons coal. The following figures are given hy King and Cawley (J. Inst. Petroleum Tech. 1936, 22, 601):

	Creosote.	Low temp.	Low temp. tar oil.	Petroleum. Diesel oil.	Gasoline produced.
Carbon % Disposable hy-	87.0	82.0	82-0	86.0	85.7
drogen % . Mol. ratio	6·7 0·92	6·9 1·01	7·9 1·17	12·2 1·7	14·3 2·0

Yields of gasoline on complete conversion are of the order of 80 to 90% hy weight or just over 100% hy volume. The octane value and other properties of the spirit depend upon the operating conditions; for further information, see Gordon (J. Inst. Fuel, 1935, 9, 69), King ibid. 1936, 9, 323), Smith (Engineer, 1936, 161,

Flotation processes for the concentration of certain minerals, particularly sulphide ores, are important and widely used. A mixture claimed to give good results is pine oil 10%, coal-tar creosote 80%, coal tar 10%—the quantity of flotation oil employed is 1-3.5% hy weight of the mineral treated. The content of phenols in the grade used should he kept constant.

Grease Making .- Attempts have heen made to employ the oils distilling above 300°C, freed from crystalline matter and having a viscosity 2-3.5° Engler at 50°C, as a home source of lubricating oils. A tendency to resinify and a high rate of viscosity change with temperature are serious disadvantages (Bruhn, Stahl u. Eisen, 1919, 39, 402, 469; Broche, Glückauf, made. A type analysis is sp.gr. 38°C. 0.995; 1932, 68, 965). These oils have been employed distillate to 205° 1.8, to 230° 18.3, to 315°

300°C. Krebs (Gas- u. Wasserfach, 1930, 73, as a sealing oil in waterless gasholders. They are also employed in the production of axle greases, the oil heing mixed with slaked lime and

> Brick Oil.—A heavy fluid creosote of sp.gr. 1.06-1.08 at 60°F.

> · Lamp (Flame) Black is manufactured by partial comhustion of the oil in a limited air supply. The yield is according to the carhon content of the oil and the design of the hurner and condensing plant; normally the figure for a mixture of naphthalene and anthracene oils is 60 to 70%. Coarse and fine grades are obtained hy fractional separation in the chambers through which the gases pass.

> LOW TEMPERATURE TAR CREOSOTE.—These oils are obtained from tars produced hy carhonising hituminous coking coals at temperatures below 600°C. The hydrocarbons present are more closely related to petroleum (paraffin and naphthene series) than to the gas-tar oil hydrocarhons, being of low specific gravity, naphthalene in recoverable quantities being absent, and the higher hoiling fractions contain paraffin wax. The unwashed oil contains a high percentage of phenolic bodies (cresols and higher homologues) and of other hodies soluble in aq. NaOH. For fuller information on constitution, see G. T. Morgan (J.S.C.I. 1928, 47, 131T) and Fuel Research Dept. papers.

> Low temperature tars may be directly hydrogenated to yield an equal volume of motor spirit or the tar may be submitted to distillation and refining processes to yield fractions suitable for employment as Diesel engine fuels, furnace fuel oils, timber-preserving oils, or for manufacture of disinfectants. Typical analyses of oils from Coalite process low temperature tar (see Bristow, J. Inst. Petroleum Tech. 1936, 22, 583) are:

> Diesel Oil.—A fraction distilling mainly between 200° and 300°C. which has been carefully washed and redistilled, and to which 2% of primer has been added, sp.gr. 0.916 at 60°F.; closed flash-point (Pensky-Martin) 166°F.; viscosity (Redwood No. 1) at 100°F. 31 seconds; coke 0.08%; cold test at 0°F. fluid; total sulphur 0.91%; spontaneous ignition temperature in oxygen 255°C. phenols nil; calorific value gross 18,450, nett 17,269 B.Th.U. per lh.

> Fuel Oil.—(a) Sp.gr. 0.983 at 60°F.; water 0.3%,; open flash-point (Pensky-Martin) 212°F.; fire-point (Pensky-Martin) 255°F.; viscosity (Redwood No. 1) at 70°F. 81 seconds, 100°F. 50 seconds; initial and final hoiling-points respectively 207° and 400°C. (91.5%); orude phenols 2.5%; calorific value 18,275 B.Th.U. per lh.; sulphur 0.78%.

> (b) An unwashed grade has sp.gr. 1.0085 at 60°F.; closed flash-point (Pensky-Martin) 186°F.; viscosity (Redwood No. 1) at 60°F. 53.5 seconds; distillate to 300°C. 72%: crude phenols 50%; calorific value 16,848 B.Th.U. per lb.;

> Timber-Preserving Oil.—A fraction to comply with B.S. 144/1936, has heen placed on the market and detailed investigations of the wood preserving properties of this oil are heing

sulphur 0.70%.

653%; phenols and compounds soluble in aq. [NaOH in distillate to 315°C., 46%

Oil for Disinfectant Manufacture.-Sp gr. 0 978; phenols, etc., soluble in aq. NaOH, 44%; distils mainly between 180° and 350°C.

BLAST FURNACE CREOSOTE resembles a low temperature tar, the specific gravity of the hydrocarbons being low, and little or no naphthalene being present; the proportion of phenols is high (25-35%). The phenols consist of cresols and higher homologues, and creosol and other derivatives of dihydroxy phenols are present, and the mixture has a high germicidal value. The product has a strong characteristic odour, resembling that of wood creesote.
Formerly produced in large quantities in
Scotland, it was employed for disinfectant manufacture and for erecoting timber

washed oil used for benzole recovery had ap gr 15 0 96; phenols 3 to 5%; distillation test, start 220°, 50% at 275°, and 70 to 75% at 300°C.; sp heat 0 42 to 0 44 cal /g /°C H. Moore quotes the following analysis: sp gr 4 0 903, distillation start 185°, 20% 200°, 82% 225° Calorific value gross 17,986, nett 17,286 B Th U per lb; viscouty (Redwood) 57 at 70°F, phenols 23%, ultimate analysis, C 82 8, H 9 9, O+N 7 22, S 0 28% Production has now practically ceased, metallurgical coke now

replacing coal for reduction purposes. by the absence of phenols It is usually rich in paraffine, is of low specific gravity, and may contain naphthelene and anthracene.

F. M. P

CRESATIN, m cresyl acctate CRESOL, C.H. (CH.)OH. The three 130. meric cresols exist in coal tar (Williamson and Fairlie, Annalen, 1854, 92, 319, 1hle, J. pr. Chem. [11], 14, 442, Tiemann and Schotten, Ber. 1878, 11, 767, 783) in approximately the following ratio; orthocresol, 35 to 40, metacresol, 40; and paracresol, 25 The erude carbolic and obtained by washing the light oil and carbolic col fractione from the primary distillation of coal tar (cf. Coal Tan, Carrotto Acid) with aq NaOH and decomposing the carbolate so chtained with CO, contains phenol in association with the three cresols and xylenels On fractionation in a still heated by ateum coils operated under vacuum and fitted with a highly efficient column, there are obtained a phenol fraction -an intermediate fraction which on cooling yields further crude phenol and a liquid of high orthocresol content-a crude orthocresol-a second intermediate fraction containing a roix. ture of the isomeric cresols-and finally a crude meta para fraction.

The two fractions of high orthogresol content are refractionated, yielding technical o cresol, which is further purified by fractional crystallisa tion. The crude meta para fraction is fraction ally redistilled to yield a product containing at least 50% metacresol (cf Cresvic Acro) or if the isomers are to be separated, a process impossible by distillation, a close-cut fraction containing 58 to 60% meta., 42 to 40% pera adds petrol (bp. 120-160°) with agitation, and cresol is collected and treated by one of the cools to obtain the m cresol compound which is following methods,

The general method is based on the separatum of the corresponding sulphonic acids. The meta para mixture is heated to 100° with four time ite weight of cone. H2SO, or the mixture is treated without external heating with three times its weight of furning sulphuric acid (200) SOa). Sufficient water is added to the charge to reduce the boling point on the passage of superheated eteam to 116-120°. The meta cresol aulphonate hydrolyses and meresol distils over. The para-acid is finally decomposed by superheated steam at 140° (Rascha, B.P. 25269)1899, G.P. 114975; Brickner, Z. anger Chem. 1928, 41, 1043, 1062) Alternstitely, Braschig proposed to separate the sulphonic and by taking advantage of the higher solubility of the m cresol derivative in sulphune acid, the paraeresol aulphonic acid being allowed to crystal lise out by prolonged standing, the meia cresol aulphonic acid being obtained from the mother liquors (B P. 18334/1899), or a soluble sodium salt auch as the sulphate, may be added in cufficient quantity to form the sodium salt of p cresol sulphonic acid, which crystallises out The separated sulphonic acids are bydrolysed by steam. If insufficient H.SO, is used, the m cresol is preferentially attacked, and the unconverted p-cresol can be extracted by bea zene (Hofmann-Larocho, F P. 434534, of also Schulke and Mayr Terrisse, O P. 281054) (G P. 268780) effect preferential aulphonation WATER GAS TAR CREOSOTE is characterised of the m-creacl by employment of H.SO, of 80 to 90% strength, and obtain a product containing 90% m and 10% p eread, which is further purified by readphonetion with in aufficiency of acid or by freezing (Elger, U.S.P. 1015616). In G.P. 148703, sodium hydrogen sulphate is employed as the agent preferentially

sulphonating m cresol. It has also been suggested to exparate the meta- and para-cresols by means of their banom salts (G.P. 53307, 152652), or their calcium

salte (G P. 152652) Rutgers (G P. 137584, 141421) treats the mexture of isomers with anhydrous oxalic and or an anhydrous acid oxalate at 100°C. By thu means the oxalic ester of p-cresol alone is formed and separates out; it is filtered off and decomposed by water into p-cresol and exale acid, the m cresol is obtained from the mother liquors According to Darzens (Compt. rend 1931, 192, 1657; B P. 107961) the p cresol forms complexes that are true addition compounds of I mol. cresol with I and 2 mols, respectively of ovatic acid, formed without climination of water. Darkens separates in cresol by means of the complex formed by the combination of ? mols m-eresol with 5 mols, anhydrous sodium acctate in the presence of an organic solvent. The complex is separated, washed, and decom posed with water.

Schering-Kahlbaum A G. (B.P. 297083) effect separation by means of the urea addition compound of m cresol. Desseigne (Mém. Poudres, 1934-35, 26, 134-157) dissolves 1.1 to 1.5 mols urea per mol m cresol in the 60:40 meta-para mixture at a temperature not exceeding 120 adds petrol (b p. 120-160°) with agitation, and separated and decomposed by water at 50°. The para-compound is obtained from the residues by means of the oxalic acid compound (see also Bentley and Catlow, B.P. 374010).

orthoCresol (2-hydroxytoluene) forms colourless crystals or a white crystalline mass which discolours on exposure to light, crystalllising-point 30.45° (Dawson and Mountford, J.C.S. 1918, 113, 925); 30·75° (Bridgman Physical Rev. 1914, [2], 3, 184); boiling-point at 760 mm. 191°C., 600 mm. 182°, 400 mm. 168°, 163 mm. 140°, 76 mm. 120°, 49 mm. 110°, (Brückner, Z. anal. Chem. 1928, 75, 289); d_4^{20} 1.0465, d_{40}^{40} 1.0290; soluble to 3.0 vol. per cent. in water at 40°; conductivity 0.127×10^{-8} at 25°; n_D^{40} 15372; specific heat 0-20°, 0.499. Forms dibromo- and dinitro-derivatives. At low temperatures yields additive compounds with pyridine and naphthalene. Standard commercial grade has crystallising-point 30·3-31·0° (B.S. 522, 1938).

metaCresol (3-hydroxytoluene) is normally a colourless, hygroscopic liquid when fresh, colourless, hygroscopic liquid when fresh, becoming slightly brown on keeping; crystallising-point 11.8° (Darzens, Compt. rend. 1931, 192, 1657); boiling-point 760 mm. 202°, 600 mm. 193°, 305 mm. 170°, 110 mm. 140°, 48 mm. 120°, 19 mm. 100° (Darzens, *ibid.*) d_4^{20} 1.0336, d_{40}^{40} 1.0184; soluble to 2.5 vol. per cent. in water at 40°C.; conductivity 1.397× 10^{-8} at 25°; n_D^{40} 1.5332, n_D^{22} 1.5352; specific heat 0-20°, 0.479; mol. heat of evaporation 10,860 cal. Forms tribromo- and trinitroderivatives. Standard commercial grade has n_D^{20} 1.037-1.040: crystallising-point not sp.gr. 15.5 1.037-1.040; crystallising-point not lower than 10.5° (B.S. 522, 1938).

paraCresol (4-hydroxytoluene) forms hygroscopic colourless prismatic crystals which discolour on keeping; crystallising-point 36-0° (Fox and Barker, J.S.C.I. 1918, 37, 268T); boiling-point at 760 mm. 202-3°, 500 mm. 187°, 217 mm. 160°, 106 mm. 140°, 47 mm. 120°; d²⁰ 1-0347, d⁴⁰ 1-0186; soluble to 2·3 vol. per cent. in water at 40°C.; $n_{\rm D}^{40}$ 1.5319 (Gibbs, J. Amer. Chem. Soc. 1927, 49, 839). Forms dibromo- and dinitro-derivatives. At low temperatures yields additive compounds pyridine and naphthalene. Standard mercial grade has crystallising-point not lower than 34° (B.S. 522, 1938).

For analytical methods, see CRESYLIC ACID. Employment-The main market demand for the cresols is for mixtures of the isomers of controlled composition particularly with respect to the content of the ortho- or meta-cresols, or for the general mixture sold as cresylic acid (q.v.). There is, however, a growing demand for the pure isomers as raw materials for the production of derivatives required by the organic chemical industry. Some of the more important uses are as follows: o-cresol is used to a limited and unimportant extent in the dycstuffs industry, the ammonium and potassium salts of dinitro-o cresol were previously employed under the name "Victoria Yellow"; p-nitro-o-chlor-o-cresol is a basic material for the

yield of m-cresol is given as 83% and the pure is employed in manufacture of cotton and azo dyes. The use of the potassium salt of dinitroo-cresol as an ingredient of plant washes has been patented; o-cresyl phosphate is a starting material for coumarin (Agfa, G.P. 246871, F. Raschig, G.P. 223684), and finds increasing employment as a plasticiser and fire retarder for nitro cellulose and celluloid. The resin from pure o-cresol and formaldehyde is odourless and has been patented as a shellac substitute.

Trinitro-m-cresol (cresylite) has been employed as an explosive. m-Cresol is the starting-point for Musk Ambrette and for synthetic thymol.

p-Cresyl methyl ether is employed in perfumery in the preparation of synthetic ylangylang oil (Schimmel A.-G., G.P. 142859, also Perf. & Essent. Oil Rec. 1923, 14, 398), and the amino derivative of this compound in the production of azo dyes. Another important derivative of p-cresol is anisic acid.

Under controlled oxidising conditions (copper oxide and caustic soda) the cresols yield the corresponding hydroxybenzoic acids and by the action of carbon dioxide on the sodium salts

the cresotinic acids (q.v.) are formed.

In general, the toxicity decreases and the germicidal value of the phenol homologues increases as the alkyl group increases in complexity (Coulthard, Marshall and Pyman, J.C.S. 1930, 280). The phenol coefficient of the cresols is $2 \cdot 2 \pm 0 \cdot 2$. A number of derivatives of m-cresol of high germicidal value have been prepared, e.g. amyl-m-cresol, phenol coefficient 200-300; 5-chloro-m-cresol, phenol coefficient in aqueous solution 133, in soap solution 24, in caustic soda solution 8 (Rapps, J.S.C.I. 1933, 52, 175T).

F. M. P. IN PER-CRESOL COMPOUNDS FUMERY. A number of cresol compounds have recently come into vogue as synthetic perfume materials. Of these the following are the most important: paracresyl methyl ether, C, H, OMe, has a well-defined odour of wallthe butyl ether is similar, but has a different unance." The phenyl ether, C₇H₇OPh, has a very powerful odour of the hyacinth-rose type, it has the following characters: sp.gr. about 1.063, ref. ind. 1.5720. p-Cresyl acetate, C₆H₄Me·OAc, is a synthetic perfume having a very penetrating odour of narcissus, and is used in many perfumes of the lily, lilae and honey-suckle types, it has sp.gr. 1.055, rcf. ind. 1.5025, b.p. 209°. p-Cresyl phenylacetate, C₆H₅·CH₂·COO·C₇H₇, is used to a considerable extent as a narcissus perfume. It is a crystalline substance melting at 75°.

E. J. P. CRESOTIC ACID. The ten isomeric acids (empirical hydroxy-toluic formula, C₈H₈O₃) are sometimes called cresotic acids, a term which indicates their relationship with the cresols. In American terminology the name is applicable to all ten isomers, e.g. 4:3-crcsotic acid is 4-hydroxy-3-methyl-benzoic acid. In the German system of nomenclature the name "Kresotinsäure" is restricted to the following production of sulphur dyes and o-cresotinic acid three compounds, which are obtainable from o-,

m., and p-cresol respectively by the action of more than traces of sulphuretted hydrogen, and carbon dioxide, sometimes under pressure, in the presence of elkalı (the Kolhe synthesis):

orthoCresotic acid, or 2 hydroxy-m-toluic acid, m p. 165°:

(For manufacture, see G.P. 138563, 624318.) metaCresotic coid, or 3 hydroxy-p-toluic acid, m.p. 177°:

paraCresotic acid, or 4-by droxy-m-toluic scid, m.p. 150°:

(For manufacture, see G.P 138563)

They are homologues of salicyho acid which they resemble in general properties. Some derivatives have been reported to have medicinal value.

CRESOTINIC ACID v. CRESOTTO ACID. ETHER CRESYL ETHER (ds cresyl oxide), (C,He),O. There are three forms, o, m, and p. The di-ortho- compound boils at 275° and is used in the preparation of artificial geranium

perfume CRESYLIC ACID. The mixture of nonerystallisable, higher boiling phenois obtained in the manufacture of carbolic acid (q.v.) from 60'a carbolic acid is, after redistillation, known commercially as cresylic acid or liquid carbone acid. When a good market demand exists, the creosote oil fractions distilling after the carboho oil fraction (v. Coal Tan) may also be washed with caustic sods solution for the recovery of the phenol contents. (For recovery processes from emmoniecal liquor, see Carbotto Acid). The product is a clear liquid varying in colour from white to dark reddish or yellowish brown, and containing 95 to 100% phenols according to bydrolysis by steam of the sulphonic soil derivagrade, the chief impurities being neutral oils tives may also be employed. The phenol mr (nephtbalene and methylnephtbalene), pyridine bases, and water. The normal refined product base, and water. The normal refined product consists mobile antility of creecis and/or sylends of the consist mobile and may contain phenol in relatively small amounts. Formerly crespite acid contained, as a cid switch are more stable of higher temperature, and any cresolain the proportions they occurred in the carebon oi, i. e. 35: 40:25, but the development of the synthetic resin (phenols formalds, high) industry baselet to a demand for product of closely standardized ortho- and/or material and the composition. The specific gravity may be within the limits 1-90 and 1-905, and the distillation renge may lie within the limits 1-90 and 220°C. The acid should be sweet smelling, contains of characterise the phenols.

react neutral to moistened htmue paper. The "British Pharmocopoeia" 1932 quelty has ap gr. 186 1-035-1-050; on distillation not more than 2% passes over below 188° and not less than 80% distils between 195° and 205°. The hydrocarbon content is below 05%, base calculated as C₅H₅N below 0 1%, readus on evaporation below 0 1%. The acid is almost completely soluble in 50 parts of water, the solution baying a pungent taste, and freely in alcobol (90%), other, chloroform, light petroleum, glycerine, and in fixed and volatile oils. The purer grades dissolve in dilute caustic soda to give a cleor solution or one from which only a few flakes of nephthalene separate en standing. B.S.S. 515, 1938, recognises fre grades, the purest grade containing not more than 0 5% impurities (water, pyridine bases, and neutral oils), and the lowest grade up to 5% impurities, of which not more than 2.5% may be neutral oils. The standard cresylic acid of high metacresol content is a colourless to pale straw liquid and has apgr. 185 1 035-1 040, metacresol content 50-55%, water content below 0.5%, neutral ollsand bases each separately below 0.1% and distils 92% between 199° and 204°C. (cf. B.S. 521, 1938, and German and Japanese Pharmacopæiss). The corresponding product of high orthocresol content contains not less than 45% orthocresol, has spgr

product (cf. also CRESOLS) . Cresylio soid is highly poisonous. The phenol coefficient as determined by the Rideal-Walker method lies between the limits 2 and 5, depending upon the proportion of higher phenols present. Cresylic acid from low temperature tars and some vertical tara tends to give a pink solution in alkahe, and a number of methods have been patented for overcoming this dis-

1-045-1 050, distils not less than 90% between

192° and 200°, and conforms to the impunty

limits quoted above for the high metecresol

advantage (B.P. 397847, U.S.P. 1822842). Methods of Analysis.—The cresols and homologues may be identified by means of the corresponding aryloxyscetic acids formed by heating the sodium salt of the phenol with chloracette acid (Fischer, Ges. Abh. Kennt. Kohle, 1917, 2, 236; Bruckner, Z. angew. Chem. 1928, 41. 1043, 1062). A method besed on the fractional ture is well shaken with an equal part of 184 H.SO, and then heeted to 103° for 3 The acid should be sweet smelling, contain not inrethanes to characterise the phenols.

the method of Potter and Williams (J.S.C.I. 1932, 51, 59), which is based on the property of o-cresol of forming a crystalline compound of definite crystallising point with cincole. A mixture of 8-40 g. of the dry sample, which must contain more than 30% orthocresol, and 12.0 g. pure cincole is prepared and the approximate then warmed to a temperature approximately 5°C. above the crystallising point until all but the last traces of crystal are melted, and the crystallising point determined by the usual method. The percentage of o-cresol is ascertained by reference to the following table, which is based on the use of cineole of crystallising point 1.2°C.:

Crystallising point of o-cresol-cineole complex. °C. o-Cresol in sample. 100 56.4 53.0 90 80 49.6 70 45.8 60 41.5 50 36.7 31.0 40

Precautions are necessary to ensure absence of

If the preliminary test shows that the o-cresol content is lower than 30%, the test is repeated using a mixture of equal parts of the sample and of pure o-cresol. The cincole and the sample tested must be quite dry. The test gives reliable results in the presence of phonol, m-, p-cresols and xylenols (J.S.C.I. 1938, 57, 212).

metaCresol is determined quantitatively by the method of Raschig (Z. angew. Chem. 1900, 13, 759) which depends upon the formation of the trinitro derivative when m-cresol is treated at hoiling temperature with nitric acid in excess. while the o- and p-crosols under these conditions are completely oxidised to oxalic acid. The sample should not contain more than 1% of water, 5% of phenol or 10% of xylenols. The m-cresol content should be between 45 and 80%; 10 g. of the sample are weighed into a 50 c.c. flask and mixed with 15 c.c. 96% sulphuric acid. After being heated for an hour at 95-100° the contents are poured into a wide-necked litre flask and rapidly cooled under the tap with a circular motion to spread the compound over the sides of the flask. 90 c.c. of dilute nitric acid (sp.gr. 1.38-1.40) are taken in a cylinder and employed in small quantities to rinse the sulphonic acid from the small flask into the litre flask. The flask is then shaken at arms' length in a fume cupboard until the sulphonic acid is completely dissolved. Gentle agitation by swirling is continued until the reaction commences as shown hy the copious evolution of brown fumes. The flask is allowed to stand for 20 minutes on a plate of insulating material and the contents are then poured into a porcelain dish containing 40 c.c. water, the flask being thoroughly rinsed out, using in all 40 c.c. water. The dish and its contents are allowed to stand 2 hours; the solid mass of trinitro-mcresol is then reduced by means of a pestle to a coarse powder and transferred completely to a

orthoCresol is quantitatively determined by tared Gooch crucible. For the transference of the remainder of the solid and the washing of the powder, a total of 100 c.c. water delivered from a fine jet wash bottle is employed. The crucible and contents are dried to constant weight at 95° to 100°C. Under these condition I g. of m-cresol yields 1.75 g. of trinitro-m-cresol. The powder should remain solid at the temperature crystallising point determined. The mixture is of drying indicating freedom from excess xylenols; otherwise the test should be repeated with the addition of a known weight of pure m-cresol to the original sample. The general conditions and, in particular, the quantity of nitric acid, must be strictly adhered to in order to prevent the development of dangerous conditions (see "Standard Methods for Testing Tar and its Products," 2nd ed.).

Phenol in cresylic acid is determined by Chapin's method (Ind. Eng. Chem. 1920, 12, 771), which is based on the fact that formaldehyde quantitatively bleaches the colour developed by treating phenol with Millon's reagent (an aqueous solution of neutralised mercury nitrate), whereas the colour developed by higher phenols with the reagent remains unchanged. See also qualitative and quantitative tests given

in article on CARBOLIC ACID.

Uses.—The total production of cresylic acid in 1935 in Great Britain was of the order of 5 million gallons, of which about 35% was exported. The U.S.A. take a special grade which on distillation yields not more than 5% phenols at 190° and not more than 75% at 215°C.; this grade is admitted duty free, the British export figure to U.S.A. in 1935 being 789,158 gallons. Chile takes 300,000 gallons cresylic acid for employment in the froth flotation process for the concentration of minerals.

The main home markets are for:

1. Synthetic resins of the phenol-formaldehyde type—the demand is mainly for a product of m-cresol content standardised within ±1½% of an agreed figure, normally lying between 50 and 58%. Cresylic acid is employed in place of phenol by reason of lower cost, consistent with good electrical and chemical resistance of the resin. orthoCresol tends to retard, whilst metacresol assists resin formation.

2. Disinfectants.—This 456,000 gallons in 1934. absorhed marketThis figure includes the acid used for lysol, emulsifying disinfectants, which give a milky emulsion with water, and which are employed as sheep dips, veterinary fluids, or for household disinfectants, soaps,

and disinfectant powders.

3. Froth flotation of minerals. 4. Synthetic tannins, tricresyl phosphate (a lacquer and varnish plasticiser and fire retarder for nitrocellulose), methyl cyclo-hexanol (solvent, plasticiser, and wetting agent for textiles), explosives.

5. Inhibitors in production of motor fuels

(high orthocontent preferred).

6. Lubricating oil refining by solvent methods (Chem. Met. Eng. 1935, 42, 246; Oil and Gas J. 1935, 33, (45), 86; 33, (51), 16).

7. Pure oresols for manufacture of derivatives required hy the dyestuffs and perfumery

CRESYLITE. An explosive mainly of 2.4 6-transtrometacresol which has been used in France for filling projectiles A mixture of this explosive with 40% of piene acid made by melting the two components under water has also been used for the same purpose under the name of Cresylite 60/40 or Cresylita

Manufacture -Trinitrocresol can be mann factured by a similar process to that used for pieric acid. In order to obtain a high yield it is (or metacristobalite, the cubic, optically isotropic necessary that the cresol used should consist modification) at 1470°. On cooling, the latter essentially of the mets modification, since the other isomers are largely destroyed during the nitration process with considerable loss of nitric

According to Wride (Arms and Explosives, 1920, 28, 47) 28 kg. of cresol are mixed with 120 kg of sulphuric acid (92%) in a sulphona tor and stirred for one hour, the nuxture attaining a temperature of about 60°C sulphonated cresol is then run into a nitrating pot similar to that used for pieric acid, contsining 98 kg, of Chili nitrate and 170 kg of nitric acid (38%) The whole of the sulphonated cresol is added in 4 hours. During the first quarter of an hour air agitation is employed to ensure complete mixing of the materials. At the end of the 4 hours air is again blown in for half an hour, after which about 80 htrea of cold water are carefully added, air egitation atill being used After atanding for about 8 hours, the mother liquid is removed and the

explosive washed and dried
Yield.—Theoretically, 100 parts of cresol
should yield 225 parts of trimtrocresol, but
by the above process the maximum yield is about

150% of the raw material

Properties - Commercial trinitrocresol is a yellow aubstance consisting chiefly of trinitro melacresol with a small percentage of other isomers. It is considerably less soluble than piorio acid in water, 1 part of the explosive dissolving in 450 parts of water at 20°C, but is readily soluble in alcohol and ether.

Puro 2:4.6 trimitrocresol melts at 109 5°C., but the melting point of the commercial product is about 100°C. It has an acid reaction and forms salts analogous to the pretates. The trantrocresylates, however, are not so sensitive to shock as the picrates

Trinitrocresol is somewhat less effective than pierre acid as an explosive and requires a power-

detonation.

According to Kast, squeous auspensions of the explosive dissolve appreciable quantities of metals such as lead, iron, zinc and copper, but have no appreciable action on aluminium

CRISALBINE (May and Baker). Gold sodium thiosulphate, used in the treatment of pulmonary tuberculosis.

CRISTOBALITE. (SiO,) crystallising in the cubic system (pseudocubic). It was first recognised by G. vom Rath ess isomer, m.p. 141°, being converted into the in 1886 as regular octahedra up to 2 mm. atable trans. form, m.p. 232°, on exposure to across, associated with tridymute, in cavities in light. The latter separates from chloroform andesite from the Cerro de San Cristobal, near methyl alcohol in orange red plates and exhibits

consisting volcanic rocks in the Rhenish district, Central which has France and Cahfornia, and in certain meteorites The milk-white, translucent crystals are twinned and exhibit optical anomalies; these disappear suddenly at a temperature of 175°, the crystals being then truly cubic; on cooling, the bi-Sp.gr. 234, hardness refringence reappears. 8-7, refractive index 1-49, birefringence 0 0005.

Quartz when heated passes into 8 tridymite at 870°, and this in turn passes into & cristobalite passes into the birefringent a cristobelite at 180-270°. The refractive index of this artificial cristobalite is about 1.484, being slightly higher than that of tridymite (1 477); mp. 1710° Cubic erystals of silica have also been obtained artificially by heating an aqueous solution of colloids] ashea with hydrofluoboric acid at 200° under a pressure of 26 atmospheres. Cristo. balite has also been detected in silics bricks.

CRITH. The weight of a litre of hydrogen st 0° and 760 mm, pressure—viz. 0 0898 g. CROCEIN ACID, 2 asphthol 8-sulphone

acid. Used in making croceine acarlet and other dyes.

CROCETIN and CROCIN. Crocetin and crocin are carotenoid components of saffron (Grocus sativus). They are related in that crocin (I, R=gentiobioso residue), the major component of the saffron pigmont, is the di-gentiobiose ester of the dicarboxylic acid crocetin I, R=H) (Karrer and Miki, Helv. Chim. Acts, 1929, 12, 985, Karrer and Salomon, shid. 1933, 18, 643).

CH CH.CMe CH.CH.CH:CMe CO.R ЁН СН СМе СН.СН СН.СМе СО∙R

Crocin, C44H54O24, is readily soluble in bot water but practically insoluble in alcohol or ether. On acid bydrolysis it gives crocetin and glucose, whilst with elcoholic ammonia the dicarboxylic acid is formed together with gentiobrose. Crocin is extremely sensitive to dilute aqueous potassium bydroxida giving a quantita-tive yield of erocetin (as potassium salt) With aqueous methyl alcoholic alkali, however, an extraordinarily facile esterification of the crocetin occurs with formation of the mono. and diinitiatory impulse to cause complete methyl esters

Crocetin, C20H24O4, is insoluble in water and in most organic solvents with the exception of pyridine, from which it separates in red coloured leaves, m p. 275-276°, which exhibit absorption mayima at 463 and 434 5 mm in chloroform; the potassium and sodium salts are yellow in colour.

Crocetin monomethyl ester, mp. separates as reddish-yellow plates from chloro-A native form of silea form-methyl alcohol. Two atereoisomeno dimethyl esters of crocetin are known, the labile Pachuca in Mexico. It has since been found in optical maxima at 463 and 434 5 mp in chloroform. On thermal degradation it yields 2:6-dimethylnaphthalene, toluene, m-xyleno together with the dimethyl ester of 1:4:8-trimethyloctatetraene 1:8-dicarboxylic acid (II), which is formed by the remarkable transformation of the methyl group side chains, has been confirmed by the synthesis of perhydrocrocetin (Karrer, Benz and Stoll, Helv. Chim. Acta, 1933, 16, 297). tion illustrated below:

The Structure of Crocetin .- Titration with alkali shows that crocetin is a dicarboxylic acid; on catalytic hydrogenation, crocetin dimethyl ester absorbs the equivalent of 7 mols. of hydrogen (Karrer and Salomon, Helv. Chim. Acta, 1927, 10, 397; 1928, 11, 513), from which it follows that crocetin is acyclic.

Since oxidation with chromic acid proves the presence of 4 extracyclic methyl groups, the 7 ethylenic linkages must be disposed between 14 carbon atoms. On this evidence a satisfactory formulation (I) for crocetin was advanced. | allocated to crocetin:

Again the degradation of perhydrocrocetin to the diketone, C₁₈H₃₄O₂ (III):

HO₂C·CHMe—X—CHMe·CO₂H

$$\downarrow \text{Br}; \text{KOH}$$

$$HO_2\text{C·CMe(HO)·X·CMe(OH)CO}_2\text{H}$$

$$\text{MeMgl} \qquad \downarrow \text{on acid chloride}$$

$$HO·Me_2\text{C·CMe(OH)·X·CMe(OH)·CMe}_2\cdot\text{OH}$$

$$\downarrow \text{Pb(OAc)}_4$$

$$\text{Me·CO—X—CO·Me}$$

and the synthesis of this diketone (6:11-dimethylhexadecane-2:15-dione) (Karrer and Lee, Helv. Chim. Acta, 1934, 17, 545) afford a further confirmation of the correctness of the structure

$$\begin{split} \text{EtO}_2\text{C}\cdot\text{CH}_2\cdot\text{CHMe}\cdot\text{CH}_2\cdot\text{CO}_2\text{Et} &\overset{\text{Na}\,;\; \text{EtOH}}{\longrightarrow} \quad \text{OH}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{CHMe}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{CH} \\ &\overset{\text{Na}\,;\; \text{EtI}}{\longrightarrow} \quad \text{EtO}\cdot\text{CH}_2\cdot\text{CHMe}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{OH} \\ &\overset{\text{PBr}_3\,;\; \text{Na}}{\longrightarrow} \quad \text{CH}_2(\text{OEt})\cdot\text{CH}_2\cdot\text{CHMe}\cdot[\text{CH}_2]_4\cdot\text{CHMe}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{OEt} \\ &\overset{\text{HBr}\,;\; \text{CHNa}(\text{CO}_2\text{Et})_2}{\longrightarrow} \quad \text{HO}_2\text{C}\cdot[\text{CH}_2]_3\cdot\text{CHMe}\cdot[\text{CH}_2]_4\cdot\text{CHMe}\cdot[\text{CH}_2]_3\cdot\text{CO}_2\text{H} \\ &\overset{\text{MeZnl on}}{\longrightarrow} \quad \text{Me·CO}\cdot[\text{CH}_2]_3\cdot\text{CHMe}\cdot[\text{CH}_2]_4\cdot\text{CHMe}\cdot[\text{CH}_2]_3\cdot\text{CO}\cdot\text{Me} \\ &\overset{\text{III.}}{\longrightarrow} \end{split}$$

In addition to crocin, crocetin, carotene, lycopene, and zeaxanthin, saffron also contains a colourless glycoside, picrocrocin (IV), the presence of which led Kuhn and Winterstein (Ber. 1934, 67, [B], 344) to suggest that crocetin is evolved by a phyto-chemical oxidation of a xanthophyll according to the following scheme:

CROCIDOLITE. (Ger. Krokydolith.) mineral of the soda-amphibole group, consisting of sodium and iron (ferric and ferrous) silicate, NaFe''(SiO₃)₂·Fe''SiO₃. It is known only in a finely fibrous form, its name, in fact, being from κροκύς, κροκύδος, wool. In the trado it is known as blue asbestos, and it is worked for the same purposes as the other varieties of asbestos (q.v.). It is, however, more readily fusible (to a black magnetic glass) than are tremolite-asbestos and serpentine-asbestos. The mineral has been long known from the Asbestos Mountains near the Orange River in Griqualand West, South Africa, where it occurs abundantly as veins, 1 or 2 inches in thickness, in jasperschists. The closely aggregated parallel fibres are arranged perpendicularly or nearly so to the walls of the veins; and the material is extracted as slabs. It is now extensively mined for commercial purposes (A. L. Hall, Asbestos in the Union of South Africa, Mem. Geol. Survey South Africa, 2nd ed., 1930, no. 12; M. A. Peacock, Amer. Min. 1928, 13, 241).

I, M, H. and F. S. S. crooidolite is especially liable to decomposition

form limonite, while the silica is set free. There all the quantitative information on digestibility or, in other words, a pseudomorph of quartz balancing of rations. For such practical purand limonite after crocidolite. This material is eoloured a rich golden yellow, and displays a silky lustre, and being at the same time very hard, it is well adapted for oroamental purposea. When cut and polished with a convex aurface, it displays a cat's eye effect, and it is extensively used for making knohs, umbrella handles, beads, etc. Such material is known as tiger's eye, though, unfortunately, in the trade the name crocadolite is very frequently misapplied to it.

The name pseudo-crocadolite would be more appropriate. In some atones there has been allenfication without oxidation, and the indigehlue colour of the original crocidohte has been preserved: material of this kind is known as hawl's eye, The atones are sometimes atsmed artificially with dyes. See also ASBESTOS. L. J. S

CROCIN v. CROCETIN CROCOITE or CROCOISITE. Lead chromste, PhCrO4, crystallising in the monoclinia system. The name was originally given in the French form crocosse by F. S. Beudant in 1832, from sposes, caffron , this was later changed to crocouste, and afterwards to the more correct form crocoste This minars lis the only chromate of any importance found in nature, and in it the element chromium was discovered. It is found usually as crystals which are bright hyscinthred in colour with a brilliant lustre; the streak is orange yellow. On exposure to light the crystals soon deteriorate in translucency and brillianay. Sp gr. 60, hardness 21-3. Crocoste is a mineral of secondary origin in veins of lead ore end is often essociated with gold. It is found at Beresovsk in the Urals (where it was discovered in 1766), Congonhaa do Campo near Ouro Preto in Brazil, Luzon in the Philippinea, Umtali in Rhodesis, and as very fine crystals at Dundas in Taamania. The mineral has been used as a pigment, being identical in composition with the artificial product chromo yellow.

CROCUS OF ANTIMONY v. ANTIMONY OXYSULPRIDE.

CRONSTEDTITE, H.Fe, Fe, Si,O, 8 variety of thuringste.

CROOKESITE, A sclenida of copper, thallium (T | 16-18%) and aliver forming compact crade protein, and other soluble constituents masses with lead-grey colour and metallic lastre is commonly far from 100, and the difference Sp gr. 69, hardness 21-3. It occurs embended in calcute in the Skrikerum mine, Småland, Sweden. Although a rare mineral, it is of interest in being one of the few minerals containing thallium in considerable quantity. It was named in 1866 in honour of Sir William Crookes (1832-1919), the discoverer of thallrum.

CROPS. The composition of crops is usually expressed in terms of a conventional aeries of analyses, which, though they do not necessarily pick out sharply any particular plant conatituent, are widely used for comparative than cereals (q v.), grown in Great Britain are

when exposed to weathering; sodium is re-purposes. Many crops are grown directly or moved, and the iron oxidised and hydrated in indirectly as feed for livestock. Virtually then results a ferruginous quartx possessing the and untritre values is based on the same finely fibrous structure of the original mineral; analyses, which are also used in the proper poses the figures are sufficiently precise and are unlikely to be supplanted by any other system. For research purposes, however, the use of this system of analyses is unwise, as misleading results are apt to be abtained. The analyses commonly carried nut are as follow (see Ministry of Agrieulture, Fertilisers and Feeding Stuffs Reguls. trons, 1932, No. 658).

1. Dry Matter .- A weighed quantity of the

sample is dried at 100°.

Ether-soluble Substance (expressed often as crude fat or oil).-The material is treated in a Southet extractor with ether or light petroleum for 3-4 hnurs, and if necessary dried, ground finely with sand, and re extracted for a further period of 1 hour. The solvent is evaporated and the oil dried at 100°.

 Crude Protein (total mtrogen×6 25).-Total nitrogen is determined by some modifies.

tion of the Kjeldahl method,

4. Crude Fibre -About 2 g of the dry material, previously extracted with light petrolenm, is pleced in a conical litre fissk, 200 ml. of hot 1 25% sulphurio acid is added and brought to the boil within 1 minute Boiling is continued gently for exactly 30 minutes, the original volume being maintained and the flask being rotated every faw minutes Tha contents of the flask are then poured into hot weter in a Buchner funnel in which a filter paper is supported by a piece of sotton aloth Filtration must be effected within 10 minutes. The residuo is washed free from acid with hot water, and washed hack into the conical flask with 200 ml. of hot 1.25% sodium hydroxida. Tha contents of the flask ere again boiled gently for 30 minutes and filtered through filter peper or cloth. The residue is washed thoroughly with hot water and dilute acid, and again with water till free from acid. It may then he transferred to an ashless filter paper after further washing with alcohol and ether, and dried to constant weight. The ash may be determined by in-concration and deducted from the weight of the fibre This correction is often omitted

5. Ash .- The ash content is determined by incineration at a dull red heat preferably in a

muffle furnsce.

The eum of the analyses, ash, crude fibre, is accordingly taken and described as "soluble carbehydrates " or " nitrogen-free extractives. The fraction frequently amounts to 50% or more of the material. A partition of the mitrogenous constituents is sometimes introduced, true protein " or " pure protein " being that fraction precipitated by a protein precipitant such as copper acetate or metaphosphoric acid. The difference between crude protein and true protein is sometimes expressed as " amides."

Analyses of some of the principal crops, other

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given in Table I. It should be emphasised according to age, variety, season, and soil that the composition may vary considerably | conditions.

TABLE I. PROXIMATE COMPOSITION OF CROPS.1

Crop.		Dry matter.	Ether- soluble.	Crude protein.	Crude fibre.	Soluble carbo- hydrates.	Ash.
5 1 6						}	
Root Crops—		050		2.1	0.7	21.0	1.1
Potatoes		25.0	0.1			8.9	1.1
Mangolds		12.5	0.1	1.3	1.0		0.7
Sugar beet		23.4	0.1	1.1	1.1	20.4	٠.
Swedes	•	12.2	0.2	1.5	1.3	8.2	0.9
Turnips	•	9.2	0.2	1.2	1.1	5.9	0.8
Carrots		13.2	0.3	1.2	1.5	9.2	1.0
Parsnips		15.0	0.3	1.3	1.2	11.3	0.9
Potato haulms		23.0	1.0	2.5	6.2	10.2	3.1
Mangold leaves		11.0	0.4	2.4	1.6	4.6	2.0
Sugar beet tops		16.2	0.5	2.0	1.6	8.7	3.4
Turnip leaves	•	11.6	0.5	2.2	1.5	5.3	$2 \cdot 1$
Legumes—							
Člover (green)		18.5	0⋅8	4.4	4.3	6.9	$2 \cdot 1$
Lucerne (green)		24.0	0.4	4.1	7.2	9.9	$2 \cdot 4$
Sainfoin (green)		20.0	0.6	3.5	6.9	7.8	1.2
Tares (green)	. 1	17.5	0.5	3.2	5.1	7.2	1.5
Field beans (grain)		86.5	1.7	25.3	8-1	48.3	3.1
Field beans (straw)		81.6	1.1	8.1	36.0	31.0	5.4
Field peas (grain)		86.1	1.9	23.1	5.7	52.7	2.7
Field peas (straw)		86.3	1.6	9.0	35.5	33.7	6.6
Soya beans (grain)	•	89.2	16.9	33.9	4.8	28.9	4.7
Soya hay	•	91.6	3.8	15.8	24.3	38.8	8.9
Clover hay		83.5	2.9	13.5	24.0	37.1	6.0
	•	83.5	2.6	14.2	29.5	29.2	8.0
Lucerne hay	•	83.5	2.5	13.2	28.0	32.5	7.3
Sainfoin hay	٠	00.0	2.0	13.7	20.0	32.3	7.3
Other Crops— Kale, thousand-head		15.8	0.4	2.2	3.1	8.4	1.7
Kale, marrow-stem	٠	14.0	0.4	2.2	2.5	6.9	1.9
		14.0	0.5	2.9	2.9	7.3	1.9
Mustard	•	11.0	0.4		2.9	5.9	1.4
Cabbage	٠	11.0	U·4	1.5	2.0	9.9	1.2
Grasses—						,, ,	0.0
Rye grass (perennial)	•	24.8	0.7	2.9	7.1	11.5	2.6
Pasture grass	•	20.0	1.1	5.3	2.6	8.9	2.1
Meadow hay		85.7	2.5	9.7	26.3	41.0	6.2
Meadow hay (aftermath)		85.2	3.4	11.5	22.5	39.3	8.5

¹ Some analyses have been taken from "Rations for Livestock," Min. of Agric. Bull. 48.

This system of analysis is more satisfactory | tion to the increase in cellulose, and contains no stituents appear in the two groups "crude fibre" and "soluble carbohydrates," the latter consisting of any non-nitrogenous material which is soluble in or hydrolysed by hot 1.25% alkali and acid. The crude fibre residue is composed almost exclusively of cellulose, but even so only represents 70-80% of the cellulose of the tissue. It may be accompanied by a small but variable proportion of the lignin. The major part of the

when applied to root crops than to mature materials such as hay. The carbohydrate constituents appear in the two groups "crude hydrate" fraction of root crops and starchy grains is more or less accurately so described, that of maturer materials is misleading in that it includes lignin and polyuronide hemicelluloses which are structural constituents associated with the "fibre."

The nitrogenous constituents commonly expressed as crude protein are not all in that condition, and frequently only 60-80% may be lignin passes into solution in the hot dilute alkali and is therefore included as a "soluble carbohydrate." For this reason analyses of a maturing material do not reveal the extent of the changes undergone on lignification, since the "crude fibre" figure only increases in proporbolism. The magnitude of the error caused by investigated, tha most recent report being that calculation of tha total nitrogen as crudo protein of Davies on mangolda (J. Agric. Sci. 1928, 16, is not great.

CROPS.

ROOT CROPS -The chief differences between the various roots are in the nature and proportions of the carbohydrate reserves stored therein. In potatoes the characteristic carbohydrate is, of course, starch, which is deposited in the cells as granules with a banded atructure. Approximations to the starch content of potatoes may be made from the specific gravity of the tnber. The amount usually stated to be present is in the neighbourhood of 20% of the fresh weight, though direct determinations of starch usually give a slightly lower figure of 70-75% on the dry weight. A small amount of angar may also be present normally, and this may be increased if the potatoes have been subjected tha low temperature in storage. In sugar beet, mangolds, swedes and turnips the chief simple carbohydrata is sucrose, stored in the root, the amounts usually found descending in the order named. The concentration of sucrose in the sugar beet is very high, though variable average figure appears to be about 16% of the fresh weight. Concentrations as high as 21-22% have been found. The aucrose is always accompanied by a certain amount of reducing sugar. Traces of raffinose may be present and may

accumulate in beet sugar molassea (v. BEET). The leaves of the sugar beet are lower in sucrose than the root but may contain much higher amounts of reducing augurs. Analysis of the whole plant, and leaves and roots separately have been given by Knowles, Watkin,

and Hendry, J. Agric. Sci 1934, 24, 368: Whole plant Leaves Roots 20 26 14-21 25-10 Dry matter . .

On dry basis. 6 60 21 00 299 Aah 14 57 53 66 71-40 Total sugars as sucroae Sncrosa 48 95 3 23 69 63 Reducing sugars . 4.73 11.34

Mangolds also contain considerable onsutities of sucrose, but not at such a high concentration, 6-8% sucrosa is commonly found. The dry matter of mangolds (10-13%) is much lower than that of sugar beet, so that on the basis of dry matter the difference between sngar beet and mangolds is not great. The yield of mangold roots per acre is more than twice that of augar beet, but navertheless, the sugar beet as the more concentrated source has the advantage. The leaf/root ratios of thesa two crops are entirely different. One part of top makes on the average 0.75 parts of root of the sugar beet, and 5 parts of root of mangolds. Swedes and turnips contain less aucrose, 4-5% having been recorded.

It should be mentioned that the cell wall of these roots contains pectin in amounts higher major part being in the form of reducing sugars, than is met with in most other tissues except not as sucrose: certain fruits. Pectin from the engar beet particularly has been the subject of investigation by Ehrhen (Ehrhen and von Sommerfeld, Brochem. Z. 1926, 166, 263; Ehrhen and Sehnbert, Ber. 1929, 62, [B], 1974). The nitrogenous constituents of root crops have not been extensively Sucrosa

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LEGUMES -The importance of leguminous crops lies in their high protein content, and in a number of cases the proteins have been fractunnated and constituent amino acids determined (s. Osborne and collaborators, J. Amer. Chem.; also Davies, J. Agric. Sci. 1920, 16, 280). Detailed analyses have been made of lucerne at various atages of growth and intervals of cutting (Graber, Wis. Agric. Expt. Sta. Res. Bull. 80, 1927). Separate analyses of leaf and stem have been carried out by Woodman (Woodman and Eden, J. Agric. Sci. 1935, 25, 576) and expressed on the dry basis:

Ash Ether soluble Cruda protein Cruda fibre	:	Young leaf 12 26 3 43 30 13 12 99	Young stem 9 78 1.20 16 97 30 77	Mature leaf. 13 94 3 01 23 48 13 39	Mature stem 7 03 1 03 9 44 44 36
Soluble carl	m.				

. . 41-19 41 28 46 18 38 14

The nature of the fraction described as aduble carbohydrates has not been examined, but it is probably composed to a considerable extent of hemicelluloses and lignin together with about 10% starch. The leguminous seeds also contain starch, the granules being of characteristic aize and shape. Soys beans are unusual in containing up to 20% oil, which has found commercial uses. The ash of leguminous plants

hydrates

(other than seeds) is usually distinctly high in calcium, as will be seen in Table II. OTHER CROPS - Because of its incressing use as a winter feed kale has been the subject of investigation by Woodman (Woodman, Evans and Eden, J. Agric. Sci. 1936, 26, 212). The compositiona of the two leading varieties, marrowstem and thousand head, ora very similar:

	,		,	
Ash			Marrow- stem 11-49	Thousand- head 11:27
7.5		•		
Ether-aclubic			2 61	2 56
Ether-soluble Crude protein Crude fibre Enluble carbohydrate			14 64	15 81
Cruda fibre			16 34	21 29
Enluble carbohydrate	es,		52-72	49 07

Separata analyses of the leaf, marrow, and rind have been made in the case of marrow-stem kale .

	Leaf.	Marrow.	Rind.	
Aah	14.86	15·19	7 05	
Ether-soluble	3 3 3	1.34	0.81	
Crude protein	13 67	12 03	7 72	
Crude fibre	12 45	13 06	29 OS	
Soluble carbohydrates	55.49	58 38	55 34	

The marrow of the lale atem is rich in sugar, the

		Marrow s	tem kale.	Thousand Nov. 20.	head kale.
	Total augare	. 54-37	45 53	47-57	43 91
	Reducing	. 50 32	21-85	29 90	29 24
1	sinkara .	. 00 32	21.00	20 00	34.07

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widely according to the species, age and manurial conditions. Young spring growth is higher in protein than summer growth, unless content. Analyses of the nitrogenous constituents of grasses have been made by Fagan (Welsh Plant Breeding Station, Bull. Series H, No. 3, 1924; No. 9, 1926–27), by Woodman, and by S. J. Watson (J. Agric. Sci. 1928–1936).

Ash of Crops.—The present as true present as tr nitrogenous fertilisers are applied. Aftermath present as true protein to be more or less steady in intensively grazed pasture grass, the value being usually between 85% and 90% and rarely below 80%. Abnormally high proportions of non-protein nitrogen have occasionally been observed and have been believed to be the cause of digestive disturbances in stock.

The figure for soluble carbohydrates in grasses largely to the deficiencies of the system of in Table II.

GRASSES .- The composition of grass varies analysis, as mentioned earlier. Small amounts of fructosans have been reported in grasses, and amounts up to 30% have been found in young rye grass, disappearing as maturity is approached (Norman, Biochem. J. 1936, 30,

advances in animal nutrition and the recognition of the rôle of minor elements in development and maintenance of condition. Aspects of this question are dealt with by Sir John Orr ("Minerals in Pastures," 1929). The composition of the ash was originally studied as a result of the view that the fertiliser requirements of the plant could best be determined by a knowledge of the constituents taken away from the soil by that and hays is commonly 40-50%, but this is due particular crop. Some ash analyses are given

TABLE II. Composition of the Ash of some Crops.

			Ash on dry			As	% of as	h.		
	 		basis.	CaO	K ₂ O	Na ₂ O	MgO	P ₂ O ₅	CI	SiO ₂
Potatoes			3.99	1.53	58-27	0.06	3.48	12.86	7.27	0.58
Mangolds			6.70	3.58	35.90	19.34	2.13	9.34	3.87	0.97
Sugar beet			2.99	15.39	27.42		-	18.40	4.02	4.02
Swedes			5.58	12.12	39.17	5.24	3.02	9.83	2.48	0.72
Clover			8.15	37.37	13.10	0.58	5.10	3.75	2.26	0.54
Lucerne (leaf)			13.94	40.10	15.35	0.50		4.52	3.59	0.86
Lucerne (stem)			7.03	25.46	33.01	0.57		6.69	5.12	1.28
Beans (grain)			3.27	5.60	40.58	2.12	6.61	30.69	2.43	0.4
Beans (straw)			5.80	28.39	8.07	14.57	3.12	2.31	2.49	3.08
Peas (grain)			2.53	5.74	44.31	0.39	8.97	29.30	0.45	0.3
Peas (straw)			5.87	40.34	11.78	9.90	8.30	8.26	3.84	10.66
Lucerne hay										
Kale, thousand-head.			11.27	21.22	34.44	1.24	3.91	8.78	9.85	0.2
Kalc, marrow-stem .			11.49	21.32	33.85	2.26	3.39	7.83	11.23	0.8
Rye grass			9.59	8.02	-			6.77	6.77	45.48
Pasture grass			9.03	8.71	35.40	5.13		8.64	9.62	21.70

OTHER METHODS OF ANALYSIS.—The system of analyses of organic constituents described above is official in the sense that the evaluation of feeding-stuffs is based upon it. There is, however, a distinct movement towards analyses to over-estimate the hemicelluloses at the constitution of which determine as closely as possible each expense of the cellulose. Examples of analyses important constituent instead of indefinite by this system are given in Table III.

Table III. PROXIMATE COMPOSITION OF SOME PLANT MATERIALS (WAKSMAN AND STEVENS).

	Maize stalks.	Lucerne.	Timothy bay.	Oak leaves.
Ether and alcohol soluble fraction Hot water soluble fraction Hemicelluloses Cellulose Lignin Crude protein Ash	3·42	10·41	2·90	9.93
	28·27	17·24	16·92	10.26
	20·38	8·52	22·07	15.62
	23·05	26·71	26·09	17.18
	9·68	10·78	16·21	29.66
	2·61	8·13	3·25	3.47
	7·40	10·30	5·68	4.68

Figures for cellulose and lignin contents would has no harmful physiological action (Cherhules frequently be of value in considering the com- and Bernhard, Helv. Chim. Acta, 1932, 15, 464. position of crops and particularly in following 855, 978). developmental changes which are very imperfectly reflected in simple crude fibre determinations. Detailed studies of this nature have been carried out on a variety of materials (Norman J. Agric. Sci. 1933, 23, 216; Buston, Biochem J. 1934, 28, 1028; Phillips and Goss, J. Agrie. Res. 1935, 51, 301; Norman, Biochem. J. 1936, 30, 1354). Although, of the atructural constituenta, cellulose and lignin may be determined with some degree of accuracy, the same cannot yet be said of the hemicelluloses A laborious procedure of direct extraction and weighing has been suggested (Preece, Biochem. J. 1931, 25, 1304), and used on grasses and leaves (Buston, Brochem, J. 1935, 29, 196), but a more expeditious and accurate method of determining this important group of polyasechandes 13 required.

A. O. N.

CROTALINÆ v. BLOOD ANTICOAQULANTS. CROTINE t. CROTON OIL.

CROTON OIL (Oleum crotonis, Oleum tiglis of the Pharmacoponas; Fr husle de petits pignons d'Inde) is obtained from the seeds of the shrub or small tree Croton Tiglium L. (Fam. Euphorhiacea), which is indigenous to S. India and Ceylon but found in cultivation in other tropical regions of S. Asia and Oceania

The oil, which amounts to about 30-45% of the whole seed (equivalent to 43-63% of the kernel) is recovered commercially by espression or by extraction with solvents (alcohol, ether, etc.) from the shelled or unshelled seeds accord ing to the requirements of the various national Pharmacopoeias1; the colour of the oil, its physiological activity, and its solubility in alcohol (which depends elicify on the content of free fatty acids) depend on the method employed for its extraction

Both seeds and oil should be handled with extreme caution, as they not only possess dangerous toxic and purgative properties when taken internally, but also display a serious vesicating action when applied externally to the skin (distinction from Curcas Oil, qv). (According to Cherbuliez, the purgative effect may also follow the application of the oil to the akin.)

In addition to the vesicant end purgative principles, which pass into the expressed or extracted oil, the seed kernels also contain two tosalbumins-crotonglobulin and crotonalbumin-(which were formerly classed together under the name "crotine" (see Stillmark, under the name crotine (see Stillmark, Arb. Pharmak, Inst. Dorpat, 1889; Elfstrand, Hahil. schrift, Upsala, 1898; Kobert, Landw. Versuchs Stat. 1913, 79-80, 97)), sucrose and glycose glycosides, end about 1-4% of a glycoside, crotonoside, which has been identified as the d-riboside of 2-hydroxy-6 amino-purme (seguanine), and which (in small doses at least)

¹ Descriptions of Oleum crotoms (Oleum Tagin, which appeared in former ceitions have been omitted from the most recent revisions of the British (1932). United States (1936), and Dutch (1926) Pharma-copecias; linkments containing croton oil still appear in the B.P.C.

Croton oil is used to a limited extent as a drastie cathartic (usually in admixture with castor oil or other diluent), and more frequently as a constituent of rubefiant limiments, etc , in cases of pneumonia, sciatica, etc.

The vesicating action of the seeds and oil is associated with an amorphous material, termed "Naturgiftstoff" by Flaschentrager (Böhm and Flaschentrager, Arch. exp. Path. Pharm. 1930, 157, 115; 1c, snfra) and "Résine de croton" by Cherhuliez, Ehninger, and Bernhard (Helv. Chim. Acta, 1932, 15, 658), which can be extracted hy purely physical methods (extrac-tion with methanol) from croton oil, or in somewhat higher yield (c. 1-3%) from the seeds themselves. This material varies in its properties according to the manipulation employed for its recovery, end appears to consist of a complex mised unsaturated eater (or mixture of esters) containing some free hydrosyl (phenolic !) groups: It is extremely sensitive to alcoholysis, especially in the presence of alkalis, and it is almost certain that the glassy, toxic croton resin, recovered by even such mild methods as were used by Dunstan and Boole (Proc. Roy. Soc 1895, 58, 238), Böhnt (1915, Ic., infra), and other early investigators represented a product of the partial decomposition or modification of the original natural toxin Croton resin, like the "Naturgiftstoff" from which it is derived, displays the toxic and vesicating properties of the oil to an enhanced degree, but is very easily saponified by alkalia (see Dunstan and Boole, Ic.; Flaschantrager, GP 638004, end le., infra*), yielding a complex mixture of physiologically mactive products: these include a crystalline tricyclic phenolic compound, phorbol (C₂₀H₁₈O₂I), eontaining five hydroxyl groups and three unsaturated linkings, and about 32-35% of a complicated mixture of fatty acids, among which formic, acetic, iso hutyric, tiglie, caprylic, capric, lauric, myristic, palmitic, oleic and linohe acida have been identified (Bohm, Arch, exp Path, Pharm, 1915, 79, 138; Spies, J Amer. Chem. Soc. 1935, 57, 189; J. Org. Chem. 1937, 2, 62); higher saturated acids and more unsaturated acids

appear to be absent.

Little is known concerning the cause of the purgative action of the oil, which would appear to be attributable to some factor other than the croton resn, since Cherbuhez finds that croton oil, from which the bulk of the toxic resin has been extracted, atill retains its cathartic activity. The "crotonoleic" acid of Buchheim (Wagners Arch, Heilk, 1872, 13, 1, 20; 1873, 14, 1), and of Kobert (Chem. Ztg. 1890, 14, 656) has been shown by Dunstan and Boole to have been an impure mixture of croton resin and fatty acids. Both croton oil and croton ream

**Phorbol and the product of its berzoylation are physiologically inactive, but the third product of physiologically inactive, but the third product of physiologically inactive, but the third product stde phorbol picked in the product of picked phorbol product and product of phorbol product and product of phorbol phorbol product of phorbol, phenolic (molic) derivatives are obtained.

hydrogenation.

The following analysis of croton oil by Flaschenträger and von Wolffersdorff (Helv. Chim. Acta, 1934, 17, 1444; cf. Schmidt and Berendes, Annalen, 1877, 191, 94) indicates roughly the quantitative composition of the oil (the "percentages" are only approximate, since the item of "loss" amounts nearly to 14%): "Naturgiftstoff" (toxic resin) 3.4% (the combined acids present in this fraction are not included in the subsequent figures); oleic acid 37%; linolic acid, 19%; arachidic acid, 1.5%; stearic acid, 0.3%; palmitic acid, 0.9%; myristic acid, 7.5%; lauric, tiglic, valeric and hutyric acids, each in traces less than 0.05%; acetic acid, 0.6%; formic acid, 0.8%; acids insoluhle in light petroleum 6.9%; "undetermined acids," 0.6%; glycerol, 7.2%; unsaponifiahle matter (including a sterol, m.p. 131°C.) 0.25%; loss 13.8%. As the whole of the "Naturgiftstoff" was not removed from the oil hy the method employed, the results of the analysis are not inconsistent with the opinion of Böhm that all the lower (volatile) fatty acids, including the stable solid, unsaturated tiglic acid (CH₃·CH:C(CH₃)·COOH; m.p. 64·5°C., isomeric with angelic acid), are associated with the vesicant principle, and do not form any part of the glycerides constituting the true fatty oil of the seed. Adriaens (Congo, Dec. 1934) reports 89-90% of insoluble fatty acids in croton oil, of which 9.5-10.5% are stated to consist of saturated, and 89.5-90.5% of unsaturated acids. As much as 0.6% of tiglic acid has been isolated from crude croton oil (Geuther and Fröhlich, Z. Chem. 1870, N.F. 6, 549, 576).

Croton oil is strongly dextrorotatory, $[a]_{D}^{c,20}$ +5° to +9°; owing to its remarkable composition, which differs widely from that of all other oils, its recognition by means of the ordinary quantitative reactions is not difficult: the following figures illustrate the usual ranges: m.p. -7° to -16°C., sp.gr. ¹⁵ 0.9320-0.9501; $n_{\rm D}^{20}$ 1.4734-1.481; acid val. 2-55; saponification value 200-215; iodine value (Wijs) 102-115; Reichert-Meissl value 12-13-6 (Lewkowitsch), 6.4-8.4 (Adriaens); Polenske value 1.3 (Adriaens); true acetyl value 19.8-32.7 (Lewkowitsch), 5.6-11 (Adriaens). In admixture with other vegetable oils, croton oil may be detected hy the characteristic reddish ring which appears at the junction of the liquids when a solution of the oil in two volumes of absolute alcohol is poured onto a concentrated solution of potassium or sodium hydroxide (Comte, J. Pharm. Chim. 1916, [vii], 14, 38; the reaction is due to the toxic resin).

The oil obtained from the seeds of an allied E. African plant, Croton megalocarpus Hutch (=C. Elliottianus Pax and Eng.), and known in pharmacy as Oleum crotonis Elliottianus or Ol. Elliott., has a comparatively non-irritant purgative action, and does not appear to contain any vesicating constituent; the characteristics

lose their specific physiological action upon of the oil, viz. sp.gr. 15 0.925-0.927, saponification value 188.5-201.5, iodine value 138.6-147.5, insoluble fatty acids 94-94.8%-also differ considerably from those of croton oil (cf. Bull. Imp. Inst. 1907, 5, 237; 1915, 13, 39; 1923, 21, 206; Cash and Dilling, J. Pharm. Exp. Therap. 1914, 6, 235; Martindale and Westcott, Extra-Pharmacopæia). E. L.

> CROTON RESIN v. CROTON OIL. CROTONALBUMIN v. CROTON OIL. CROTONGLOBULIN v. CROTON OIL. CROTONITRILE v. ALLYL CROTONOLEIC ACID v. CROTON OIL. CROTONOSIDE v. CROTON OIL.

CROTONYLENE v. ACETYLENES.

CROTOPHORBOLONE v. CROTON OIL. CROWBERRY (Empetrum nigrum). leaves of the crowherry contain a wax composed of ceryl cerotate, benzoio acid, ursone, tannin, dextrose, and probably rutin. The anhydrous ursone, C₁₀H₁₆O, is identical with that obtained from the leaves of Uva Ursi. Empetrum nigrum contains no alkaloids or andromedotoxins or glucosides decomposed by emulsin (van Itallie, Pharm. Weekhlad, 1918, 55, 709).

CRUCIBLES. Crucibles may be described as hollow vessels constructed of material which can resist the action of heat and chemical attack and possess mechanical strength at high This description takes no temperatures. account of size and thus embraces types ranging from small lahoratory utensils to industrial plant; and crucibles which will give satisfactory results in service are some of the most difficult refractory articles to produce. The requirements of a good crucible may be hriefly outlined

under the following heads:

MECHANICAL.—Crucibles must be sufficiently strong to resist, without hulging, the pressure of the molten material they contain. In many cases, for example, the casting of metals, alloys and glass, they must he rigid at high temperatures, so that they can he lifted and handled hy tongs, or some gripping device, without failure. The requisite strength can be secured by using, say, a suitable clay, such as those from Stour-hridge, Derhyshire, Klingenherg and certain localities in France and Belgium; hy correct grading and proportioning of the grog and admixed materials, hy avoidance of cracks and strains during manufacture, hy careful firing, and hy attention to shape and design. As a rule, attempts to increase mechanical strength hy incorporating fibrous material, such as ashestos, in the wall, or hy hinding the wall with metal hoops, have not heen successful.

Physical.—Texture, a property which can he controlled hy the grading of the component materials, must he selected according to the purpose for which the crucible is to he used. close interlocking texture is essential for mechanical strength, for preventing permeation hy molten material, and is generally stated to favour heat conductivity. In using crucibles for casting, however, they are often completely removed from the furnace and thus subjected to an ahrupt change of temperature which renders them liable to fracture. In the case of elay materials, resistance to this is secured by the

Owing to a confusion in the nomenclature, the seeds and oil discussed in the papers quoted were incorrectly described as from C. Elliotianus Baill., which is a related Madagascan species (Priv. comm. from Imperial Institute, 1937).

use of a relatively large proportion of coarse grog and that a thin lining of chalk renders carthen penetration, so that in this, as in many other points, a compromise must be etruck. Compared with metals, the heat conductivity of refractory materials is low, though in crucibles in most cases the opposite quality would be a great advantage and it is largely for this reason that grephite and carborundum have been used extensively.

CHEMICAL,-The main chemical considerations are the relation of chemical composition to refractoriness and to the resistance to the strack of molten metals, elags, etc. The refractormess fluxes, such as alkalis, iron compounds, etc., but also on the ratio of the more infusible on the other hand, a small amount of somewhat more fusible, and by the character of the atmo-

refractormess. Resistance of the inside of a crucible to chemical attack by molten metels, elags, salts, etc, ie obviously a very complex and wide question. The general principle of using an acid (silenous) refractory in contact with an acid material, and a basic (aluminous, magnesian) with a basic, bolds good. In many cases, however, crucibles are composed largely of neutral refractories, such as graphite and carborundum, that is to eay, those which do not tend to combine with either acid or basic material, a fact which hmite the range of attack to some extent. In addition to the eclection of a cuitable crucible for a particular purpose, on the linee just indicated, increased protection is often provided by uning the crucible with a layer of come specially resistant material. Assuming that the layer adheres satisfactorily, this procedure has the advantages that opecific resistance to individual forms of chemical attack is provided and the inherent properties of the crucible, such FACTURE OF CRUCIBLES .- The following simple as a coarse texture, can be retained and con- classification, which is adequate for most acquent permeation of the wall restricted, by developing a dense and compact layer on the inside. The following quotation from Percy ("Metallurgy," 1875, p. 141) is of interest:
"Lining or Brasquing Crucibles with other than

use of a relatively large proportion or course goog and the sea of a carbon carbon carbon from this first sirve). Such an admixture, coupled with from this fixpears that at that time the sivan-exclusion of finer and intermediate grades, increases the poposity and the tendency to that that other materials could be used in special that that other materials could be used in special. cases. The term "earthen crucibles" spplied to what would nowadays be described as "clay crucibles." Also, the mention of molten htharge suggests come points with regard to the destruction of crucibles by molten material. Many metals have little effect, but various salts and slags either permeate the wall or exert a definite solvent action on it. In this connection reference may be made to the ecction on clars m Percy'a "Mctallurgy" (pp. 46-86), in which the action on clay crucibles, of clags of varying composition, is epecifically described in several depends not only upon the low content of cases. From a large number of suggestions, the following are mentioned as illustrations. For erucibles made of refractory clay and eand for exides to one another. For example, the metallurgical use, Quencau claimed the use of addition of silica to some clays may reduce Innings of refractory clay mixed with a substance the refractoriness of the mixture. Occasionally, capable of resisting the chemical operations concerned, and firmly compressing the hoing before on the other many a characteristic of the first crueble. R. H. Stone J. Amer. Ceram. Soc. 1933, 16, 95-101) considers that the presence of sodium silectic (U.S.P. 1572476; J.S.C.I. 1926, a certain amount of flux (febpar) in the Klingen-45, 330). For the "Thermit" process, Gold. berg clay accounts for its suitability for graphite schmidt recommended lining crucibles with a crucibles. Apart from chemical composition, mixture of soluble glass with magnesia and the temperature at which a crucible begins to corundum (F.P. 354597; J.S.C.I. 1905, 24, soften or melt is influenced by other factore, such [1114). Deppeler claims to make the thickness as grain eize, the finer grains appearing to be of the crucible wall sufficient to remove so much beat of the reaction that a thin layer of slag aphere in which they are heated, a reducing is formed on the inner surface ac soon as atmosphere generally tending to lower the reaction begins (USP. 1530395, JSCI. 1925, 44, 403). Crucibles may be rendered non-porous by treating with ferrous sulphate solution and firing (Mitchell, U.S.P. 804845, J.S.C.I. 1908, 27, 886) Sato suggests soaking alumina crucibles (for a Tammann furnace) in an aqueous solution of aluminum chlorids and heating to above 345°C., the temperature of decomposition of the salt (B. 1935, 804). The fracture of crucibles by expansion of anddifying metal can be prevented by hining with friable material (Colby, U.S.P. 830208; J.S.C.I. 1906, 25, 934). For preparing fused quartz, Keyes euggests hining a carbon crucible with a paste of sodium tungstate endan appropriate binder; the paste is apread over the inner surface of the crucible which is then sgnited in order to harden the lining, and this is treated with a burnishing tool or roller to give it a firm and compact surface (U.S.P. 1249637; J.S.C.I. 1918, 37, 89A).

CLASSIFICATION, COMPOSITION AND MANUpurposes, arranges crucibles under the heads: Clay Crucibles.

Carbonaceous Crucibles.

Crucibles of Miscellaneous Materials. Clay Crucibles are probably the oldest and have Carbonaccous Matter." Berther (Traité des been used very largely for the melting of Essais, Paris, 1834, i. 83) stotes that he has metala, alloye and glass. The chief varieties are occasionally lined crucibles with cilics, alumins, known under epecial names which are still magnesis or chalk, previously moutened with retained, and the following descriptions, taken water, so as to make them sufficiently cohesive, mainly from Percy's "Metallurgy," may be

country are certain varieties from Stourbridge, Stannington and Derby (v. infra). A mixture of raw ground clay and about half its weight of graded grog, made from old glass-house pots from which all adherent glass has been removed, is used. The smaller crucibles are fired in a kiln, but the large casting pots are baked immediately before use, and very hard-baked crucibles are more liable to fracture on reheating. A very plastic clay, such as Dorset ball clay, which enables a large amount of grog to be incorporated, is sometimes substituted for a part or even the whole of the pot clay.

Cornish Crucibles, largely used for copper assaying are, according to Searle, composed of 1 part of Teignmouth ball clay, 1 part of Poole ball clay, and 2 parts of St. Agnes' Beacon Sand. They can be heated suddenly, but are not refractory enough for melting steel. Crucibles of this name are now made by the Morgan

Crucible Company of Battersea.

London Crucibles are round in shape, fine grained, withstand the action of litharge fairly well, but do not stand sudden changes of temperature. The name refers actually to the shape rather than the composition or place of manufacture.

White Fluxing Pots, also made by the Morgan Crucible Company, from a highly refractory fireclay from Rouen, have a very smooth surface, resist the action of fluxes well, but crack

with sudden changes of temperature.

Hessian Crucibles, largely used for melting precious metals and chemicals, are triangular in cross-section and are of varying sizes which fit into one another to form a "nest." They are somewhat porous, rapidly attacked by litharge, and very liable to fracture with abrupt changes of temperature. According to Searle, they are made of 2 parts of Gross Almerode clay and 4 or 5 parts of Epterode sand; the clay melts at Seger cone 33 to 34 and has the approximate composition:

						%
Silica						64
Alumir	ıa					24
Loss or	ı ig	niti	on			7
Iron or	xide	e, lir	ne,	etc.		5

They are obviously acid in character and in many ways resemble Cornish crucibles.

French Crucibles are made from I part of clay (Ardennes) and 2 parts of the same clay fired as grog or chamotte. They are refractory, fine

grained, and resistant to slags.

Silica Crucibles may be made by mixing sand or powdered silica with an absorbent material, e.g. kieselguhr (Sankey, B.P. 3353, 1906; J.S.C.I. 1907, 26, 96), or with cement and water and, after drying, heating to the fusing point of silica (Reppey, U.S.P. 905295; J.S.C.I. 1909, 28, 24), or with a little lime (Berglund, F.P. 366830; J.S.C.I. 1906, 25, 1081).

The tables shown on p. 438 give the composition of some clays used for crucibles and of

various types of crucibles.

Carbonaceous Crucibles .- It has long been

quoted. The clays generally used in this tendency to break with sudden changes of temperature and increases the heat conductivity without lowering the refractoriness. Carbon is generally introduced in the form of graphite, which should have a low content of mineral matter and a suitable texture. The ash should not exceed 5 to 10% and must be infusible and have no tendency to combine with the clay to form a fusible mixture. In texture, graphites consist mainly of thin flakes or of irregularly shaped grains; the former tend to give a mixture which is "slippery," whereas the latter tend to pack closely, especially if grains vary in size. Graphite which has been imported during the past hundred years from Ceylon is particularly suitable. It contains about 95% of carbon and the grains are irregular in shape and size, ranging from minute particles up to several millimetres along their maximum dimension. During the last thirty years graphite has been imported from Madagascar and yields about 80 to 90% of carbon; the flakes are somewhat straight-edged and approach parallelograms or polygons in form (Thiessen, J. Amer. Ceram. Soc. 1919, 2, 508; Muller, Ber. deut. Keram. Ges. 1935, 16, 410).

For the manufacture of crucibles, the graphite is ground, sifted, digested with dilute sulphuric acid to remove the greater part of the iron, and mixed with water and kneaded to a stiff paste with about an equal weight of finely ground clay. The mixture is passed through a pug-mill and cut into blocks. When required for use, the blocks are again passed through the mill, and then shaped, dried, and fired in saggers. The shaping may be done by hand, or by "throwing" on a potter's wheel, or by pressing in a mould by hand or mechanical power. Various moulds have been designed for the purpose, and the selection of any one is determined by the size or other properties required in the crucible. A common form consists of a large block of wood (or iron) with a deep cavity, the curved surface of which gives the shape to the outside of the crucible. The block is known as a "flask," and is partly filled with the clay mixture into which a plug or "stopper" is forced with a rotary movement, and thus shapes the inside of the crucible, the space between the flask and the stopper being filled with clay which forms the wall of the crucible. At the bottom of the stopper is a pin, or pivot, which fits into a hole in the bottom of the flask, and the hole thus left in the base of the crucible has to be filled up very carefully when the moulded piece is withdrawn. Crucibles, like other ceramic articles, are also made in jiggers and jolleys, which shape a mass of clay into the required form on the outside and inside. Lastly, crucibles are made by the casting process, in which a slip is poured into a plaster of Paris The surfaces of the best quality crucibles are finished by rubbing with a wet sponge or leather which produces a smooth and less pervious layer on the surface.

Crucibles in which carbon is the predominating constituent have been made in various ways. Small crucibles have been made by turning out of recognised that the incorporation of carbon a solid block of gas-carbon on a lathe, and with the clay of crucibles diminishes their by moulding mixtures of anthracite, charcoal,

CRUCIBLES.

ANALYSES OF SOME CLAYS USED FOR CRUCIBLE MAKING.

	1.	п	III.	īv.	y.	, A1
SiO, Al ₂ O, MgO	79 25 13 57 0 42	48 04 34-47 0 45	45-73 34-14 0-74	48 08 36 89 tr.	65 98 23 37 0 53	49 12 35-73 0 24
CaO . K.O	0 08 1-38	0 66 1-94	0 79 0 45	0 55 1-88 2 36	0 41 0 76	0 18 1-46
Fē ₁ O ₃ Na ₁ O	_	3 05	(FeO) 1-76 (FeS ₂) 0.77	-	2 32	0 56
TIÓ _k Loss on ignition	5-11	11-15	10 17	10 87	6 84 (S) 0 05	0 11 11-95
			Moint. 4-45 Org. 070			
}	99 81	99 76	99 70	100 63	100 26	99 74

Stourbridge (Percy).

II. Stannington, near Sheffield (Bischof, "Gesammelte Analysen"),

III. Edgemont, near Sheffield (Percy).

IV. Edensor, near Derby (Percy).
V. Gross Almerode (Buchof, "Gesammelte Analysen").

VI. Ball Clay (Mellor and Heath, Trans. Ceram. Soc. 1909, 7, 151).

COMPOSITION OF CERTAIN VARIETIES OF CLAY CEUCIDLES

	1	11	111	īv	v.	vı	VII	vIII	ıx
SiO ₂ . Al ₂ O ₅ . Fe ₃ O ₂ . MgO .	64 6 34 4 1 0	723 195 39 H ₁ O I8	70 9 24 8 3 8 tr.	71 0 23 0 4 0	65-2 25 0 7 2 tr.	67 4 32 0 0 8 tr.	68 0 29 0 2 2 0 5	72 39 25 32 1 07 tr. CaO 0 38	53 92 40 57
	100 0	97.5	99.5	980	97-4	100 2	99 7	K ₂ O 1 14	

- French, made by Beaufay at Pares (Berther, quoted by Percy).
- II. French, made by Deyeux at Savignies (Berthier, quoted by Percy).

- III. Hessian (Berthier, quoted by Percy).
 IV. English ateel (Berthier, quoted by Percy).
 V. St. Etienne ateel (Berthier, quoted by Percy).
- VI. Glass pots, Nemours (Berthier, quoted by Percy).
- VII. Glass pots, Bohemia (Berthier, quoted by Percy).
- VIII. Cornish, made by Juleff (Dick, quoted by Percy).

IX. Steel crucible, clay (Brand, quoted by Percy).

Dighy recommends the following composition. Plumbago, 17 mesh granules 17 parts Plumbago, 80 mesh granules Russian clay . China stone . Б ,, Gas carbon . 2

The last three ingredients are ground and mixed with water and then the plumbago is melted under reducing conditions, and in the added and the crucibles formed, dried at 65°, case of steel, for example, may be the means of and baked (B.P. 3661, 1893).

gas carbon, etc., often with an addition of tar heat in the absence of air, whereby the carbon (Putz, D.R.P. 181979; J.S.C.I. 1907, 26, 1030). is deposited in the pores. "Salamander" crucibles, made by the Morgan Company, contain a smaller amount of carbon and resist sudden temperature changes well.

The table on the next page gives the composation of some carbonaceous crucibles.

Carbonaceous erucibles, besides being highly refractory and resistant to sudden changes of temperature, ensure that material in them is introducing more carbon (Brand, Chem. Zentr.

W. Smith (B.P. 149410, 1919; J.S.C.I. 1920, 15, 407; Dingl. poly. J. 1835, 255, 223).

38, 055A) saturates porous clay crucibles with a Although the carbon is, of course, burnt away auntable hydrocarbon, and decomposes the by duming heating in a furnesc, the more fusible

	ı.	п.	m.	IV.	. v.	VI.
Carbon SiO ₂	18·60 42·78 34·71 ————————————————————————————————————	40·43 24·63 27·89 — — — 0·27	20·0 51·4 22·0 3·5 0·2 1·8	34·50 45·10 16·65 0·95 — 2·50	25·5 50·0 20·0 1·5 0·5 3·0	39·77 38·73 16·54 1·70 0·85 0·70 — TiO ₂ 0·52 MgO 0·34 K ₂ O 1·02
	96.58	93.22	98.9	99.70	100-5	100-17

I. Steel crucible, coke clay (Brand, quoted by Percy).

II. Steel crucible, graphite (Brand, quoted by Percy).

III, IV, and V. English blacklead (Mene, quoted by Percy).

VI. Salamander (Hancock).

mineral constituents in some cases form a pro-tective glaze over the carbon particles. This tective glaze over the carbon particles. This object may also be achieved by coating the crucible with clay and borax, etc. (Woolford, B.P. 24479, 1898; J.S.C.I. 1900, 19, 52; McDonald, U.S.P. 828954; J.S.C.I. 1906, 25, 119).

Crucibles of Other Refractory Materials.—So many different refractory substances have been suggested for the manufacture of crucibles that only a few illustrative examples can be given. On account of its refractoriness, alumina has often been used. Deville (Ann. Chim. Phys. 1856, [iii], 48, 195) employed a mixture of strongly calcined and gelatinous alumina, and also in place of the former, a material made by calcining an intimate mixture of alumina and powdered marble. Generally, crucibles are made from strongly calcined alumina, or bauxite (q.v.), mixed with a refractory clay as binding agent. Caspar (U.S.P. 886111; J.S.C.I. 1908, 27, 568) shaped small pieces of amorphous aluminium hydroxide by the addition of water and then drying and firing them to a white heat, in a closed vessel. In some cases he added pure magnesia and thus typified many attempts on the same lines to form a spinel to act as a binding agent. Fused alumina, in the form of alumdum (q.v.) or corindite (q.v.) has been used for crucibles, generally with a refractory clay as binder.

The fibrous character of asbestos suggests that its use would be advantageous in strengthening the walls of vessels containing molten matter and subjected to handling. Havard ("Refractories and Furnaces," p. 228) recommends mixtures of fireclay, graphite, etc., with asbestos for crucibles. At the same time it must be borne in mind that under the action of prolonged heating, asbestos tends to disintegrate and lose

its binding power.

Carborundum crucibles are very refractory, mechanically strong, and resistant to chemical Generally they are made from a

water-glass, etc.; thus, a mixture of carborundum and ganister is pressed into shape and then heated in an electric furnace (Imray and Carborundum Co., B.P. 19493, 1902, and 3308, 1903; J.S.C.I. 1903, 22, 213, 698). A strong and very resistant material has been made with 68 parts of silicon carbide, 25 parts of flake or crystalline graphite, 7 parts of clay, and a car-bonaceous binder, such as tar, pitch or molasses, or part of the clay may be replaced by a flux such as borax (Buffalo Refractory Corp., B.P. 176436, 1920; J.S.C.I. 1922, 41, 328A).

Purified and powdered chrome iron ore, mixed with fireclay to which aluminium hydroxide has been added, may be moulded into crucibles and burned, with or without the addition, before burning, of a substance rich in carbon, such as sugar, resin or coal (Bach, B.P. 24041, 1903; J.S.C.I. 1904, 24, 64; Klein, U.S.P. 787770; J.S.C.I. 1905, 24, 547). By applying a chromium salt to the inside of a crucible and then firing, a lining of sesquioxide of chromium is produced (Placet, B.P. 28728, 1896; J.S.C.I. 1897, 16,

Lime crucibles have been made from lumps of well burnt, preferably slightly hydraulic, lime by turning on a lathe. The lime may be soaked in molten paraffin wax, resin or tar to prevent slaking (Searle, "Refractory Materials," p. 471). Larger crucibles may be made by compressing a layer of lime round the inside of a clay crucible, which has been coated inside with graphite in order to prevent combination of the silica and lime at high temperatures (Percy, "Metallurgy," p. 134).

According to Percy (op. cit. p. 135), magnesia was introduced by Caron as a basic material for making crucibles and for lining reverberatory furnaces for melting and dephosphorising steel (Compt. rend. 1866, 62, 296). The magnesite was obtained from the island of Euboea and must be very strongly calcined to prevent submixture of carborundum with a nignry tory clay to give a plastic mass (Velter, F.P. 15–30% of lightly caremou (accordingly 1917); J.S.C.I. 1907, 26, 1010; Engels, B.P. material and water to form a paste which is moulded, strongly compressed, dried, and fired to red heat. It is interesting to note that

Caudin is stated to have made crucibles from a [Kjobenhavn, 1905; C. B. Beggild, Medideletes mixture of 1 part of magrens and 1 part of om Grondand, 1912, 50, 107, and Z. Kryst. bauxite, which would preasmably yield au Mm. 1913, 51, 591; C. P. Bernard, The Cryolite artificial panch, and is indicative of several Mine at 1/cgitt, Greenland, Minnag Maganne, recent investigations on basic refractories (e.g. Chestera and Parmalee, J. Amer. Ceram. Soc. 1935, 18, 94, J.S C.I. 1935, 54, 454B). Crueibles may also be made from a mixture of calcined magnesite with from 2 to 12% of a boron compound, moistened, moulded, dried, and fired at a high temperature (Rawson and Lattlefield, B.P. 16004, 1900; J.S.C.I. 1901, 20, 992; tdyf. 1902, 52). The product is and to reason molten litharge. Cruebles of a similar mixture may be rendered porous by the addition of saw-dust (Dobell, B P. 10715, 1897; JSCI. 1893, 17, 461). According to Haward (op cit p 237) cruchles of magnesia—and other refractory outdes—are made by electrically fusing the material (Weintraub, Met. and Chem. Eng 1912, 10, 308). A layer of strongly calcined magnesie, finely ground in a tube mill, is spread on the hottom of a carbon or graphite mould. A carbon plug, covered with a thick layer of paper to allow for the aubsequent abrunkage of the magnesia, is inserted, and the spoce between it and the wall of the mould filled with powdered magnesia and the whole heated in an electric furnece to 1,500°C (U.S P 1022011, 1912). similar method is given by Kerl (" Handbuch der gesemmten Thonwearen Industrie," p 963); powdered megnesia is filled into a carbon tube, fitted between two carbon blocks, the lower of which is cut to shape the inside of the article, and on passing the current the magnesia fuses and runs down over the shape.

Small crucibles, usually for laboratory work, save been made from such refractory oxides es ave been made irom such refractory ovides os mreonia, thora, cera, etc. (see futther, Percy, 'Metallurgy'' (Fucl, etc.), 1875, pp. 110-141, B. Kerl, 'Handbuch der gesammen Thomwaren Industrie,'' 1907, pp. 332-965, Searle, 'Refractory Materials,'' 1824, pp. 646-590, Havrard, 'Refractories and Furnaces,'' 1912, pp. 228-238, C Bischof, 'Die feuerfesten Tone,'' 1923, pp. 178-193)

W C. H CRYOGENINE. Phenylsemicsrhazide

antipyrine. CRYCLITE. (Cres Krysleth) h roture fluoride of aluminium and aodium, found almost sometimes of samman and somen, the armost amount of sometimes called "Greenland spar." The name eryolite, or "ice spar" (Eisstein), refers to the general resemblance of the mineral to too (epics) and to its ready fusibility in a candle flame. The mineral was discovered by the Danes in 1794, and described by Schumacher in 1798. All the eryolite used in commerce is obtained from a single locality at lyigint (or Evigtok), on Arauk-fjord in south-west Greenland This locality was visited by Giesecke, and later by J. W. Tayler (Quart. J. Geol Soc 1856, 12, 190). The cryolito occurs as a large deposit in and by the passage of CO₀ through the solution, gness, and contains (especially near its junction Na CO₀ is obtained, while alumina separates with the enclosing rock) a number of metalle a gelations form. The decomposition of cryo minerals, such as galena, pyrites and chalybite lite may also be effected by boiling the finely

1916, 14, 202).

Cryolite is usually found as cleavable translucent masses, of snow-white colour, with s peculiar moist-looking vitreous lustre which cenders the mineral easy of recognition. H. 24 sp gr. 29-30. Crystals are rare; they are cube like in habit, and belong to the monoclinic system, with three perfect cleavages appmvi mately at right angles to one another. Although cryohite is usually white, it is sometimes coloured brown or reddish, and is said to pass down wards, in the Greenland deposit, into a black mass. The black variety loses its colour when heated. Before the blowpipe crychite fuses to a white coamel.

The composition of ervolite may be represented by the formula 3NaF-AIF, or Na, AIF,... This corresponds to AI 12 8, Na 32 8, F 54 4% An analysis of the Greenland cryolite by Chodney yielded A! 13 23, Na 32 71, Mn₂O₃ and MgO 0 83%, (Yeth Russ, Min. Ges 1846, for 1815-48, 209) Durney found in the cryolite of Miask in the Urals, Al 13 41, Na 32 31, Fe₂O₃ and Mn.O. 0 55, CaO 0 35% (Pogg Ann 1851, 83, 587) Deville found in the Greenlend cryolite 0 018% of vanadio ecid and a small proportion of phosphoric acid An analysis of cryolite from near Pike's Peak, Colorado, hy Cross and Hillebrand yielded Al 12 9, Na 32 4. F 53 55, Fe₂O₃ 0 4, Ca 0 28, H₂O 0 3% (Amer J Sci 1883, 26, 271)

Cryolite was formerly a rare mineral, hut about 1855 it was introduced into Europe commercially, and at once found numerous applica-tions in the arta. About 10,000 tons are exported annually from Greenland. By heating finely powdered cryolite with sodium or with magnesium, J. Percy and A. Dick obtained metallic aluminium (Phil Mag. 1855, 10, 364) Deville found that fused cryolite is decomposed hy the passage of an electric current, with elimination of free aluminium. At the present time, aluminium is prepared by the electrolysis of bauxite, but the addition of some cryolite is necessary to start the process and to keep it in rontamena operation (v. histantica).

A wide range of applications was suggested when Julius Thomsen showed in 1850 that cryolite is readily decomposed by heating with calcium carbonate, and that sodium aluminate may be thus obtained :

AlgFa-BNaF+BCaCO, =3Na,O Al,O3+BCaF++BCO,

If heated with quicklime, a similar decomposi-tion is effected. The sodium aluminate has been used in the manufacture of alum, and in the preparation of caustic sods for scap making. By hxiviation, the aluminate is easily dissolved out; (c. P. Johnstrup, N. Jahrb. f. Min. 1836, i, pulversed mineral with milk of lime. By adding Ref. 28; R. Baldauf, Z. pr. Geol. 1910, 18, an excess of powilered crychito to the resulting 432: O. B. Bøggild, Mineralogie Groclandica, solution, the aluminate is decomposed, the aluremains in solution:

3Na2O·Al2O3+Al2F6·6NaF $\approx 2Al_2O_3+12NaF$

Cryolite can be readily decomposed by the action of sulphuric acid, with formation of sodium sulphate, which can be easily converted into carbonate; but the process, after trial at Copenhagen, has been abandoned in consequence of the inconvenience arising from the evolution of hydrofluoric acid:

Al2F6.6NaF+6H2SO $=Al_{2}(SO_{4})_{3}+3Na_{2}SO_{4}+12HF$

The manufacture of soda from cryolite is carried on at Oersund in Denmark, and at Natrona, near Pittsburg in Pennsylvania. The mineral is also employed in the preparation of enamels on iron and for the manufacture of an opaline glass, somewhat resembling porcelain. This glass may be formed hy fusing together 100 parts of silica, 35 or 36 of cryolite, and 13 or 14 of oxide of zinc. C. Weinreb has investigated the cause of the milkiness of cryolite glass, and believes that it is due to the presence of aluminium fluoride, which, on cooling the fused mass, separates in a highly com-minuted condition, and being distributed through the glass renders it more or less opaque (Ding. Poly. J. 1885, 256, 361).

CRYOLITHIONITE. Fluoride of lithium, sodium and aluminium, Li₃Na₃Al₂F₁₂, remarkable in containing a higher percentage of lithium (5.35%) than any other mineral. occurs as large, colourless, rhombic-dodecahedra embedded in the cryolite deposit at Ivigtut in Greenland. Sp.gr. 2.77, H. $2\frac{1}{2}$ -3, n_{Na} 1.3395, m.p. 710°. The mineral dissolves in 1350 parts of water at 18°, and from this solution crystallises as cuhes.

CRYOSCOPY. As early as 1788 Blagden proved that the extent to which the freezingpoint of an aqueous solution lies below that of water—the depression of the freezing-point, as it is called—is proportional to the concentration of the dissolved substance. Nearly a century later, the researches of de Coppet and Raoult showed that when equimolecular proportions of different substances are dissolved in equal quantities of a given solvent, the solutions so obtained have, in general, the same freezingpoint. It is on these facts that the use of the cryoscopic method of determining the molecular weights of dissolved substances is based.

In comparing the effects of different substances on the freezing-point of a given solvent it is obviously convenient, in view of de Coppet's and Raoult's results, to refer these effects to 1 g.-mol. of solute and to some definite quantity of the solvent; in practice, 100 g. of the latter is taken as the standard amount. The depression of the freezing-point for such a concentrated solution as one containing I g.-mol. of solute in 100 g. of solvent cannot be determined directly; it is calculated from the depression ohserved for a dilute solution, on the hasis of the

mina being precipitated, while sodium fluoride | centration. The depression, therefore, for a solution containing I g.-mol. of non-electrolyte in 100 g. of water is a constant, independent of the particular solute; the average value of the cryoscopic constant for water is 18.5°.

For each solvent, a characteristic figure can he similarly deduced from experimental data. The evaluation of this constant—the molecular depression of the freezing point—is effected by substituting experimental values in the formula.

 $K = \frac{M}{g} \cdot \Delta t$, where Δt is the depression of freezingpoint observed for a solution containing g grams of solute in 100 g. of solvent, and M is

the molecular weight of the solute.

It has been shown by van't Hoff that, on thermodynamical grounds, the molecular depression of the freezing-point for any solvent is quantitatively related to the temperature of freezing and to the latent heat of fusion. The

relationship in question is $K = \frac{0.0198T^2}{2}$, where

T is the freezing-point of the solvent on the absolute scale, and w is the latent heat of fusion per g. of the solvent. The following table shows how far the values of the molecular depression calculated for various solvents by van't Hoff's formula, are in agreement with those deduced empirically:

Solvent.		K (found).	K (calcu- lated).	
Water Acetic acid Phenol Benzene Naphthalene . Camphor Camphorquinone	:	:	18·5 39 53 51·2 69 400 457	18·6 38·2 50·5 50·7 69·5

The van't Hoff formula assumes that the solution is an "ideal" one and that pure solvent only crystallises from it. Should the latter condition not obtain the whole method hreaks down, the depressions observed heing always too low. An examination of the van't Hoff formula was made for a number of solutions by Hückel, Kumetat and Severin (Annalen, 1935, 518, 184) and the abnormalities were shown to he due to the formation of mixed crystals of solvent and solute. Attempts have been made to connect the constant with other properties of the solvent and have been successful in certain cases. Thus J. F. Durand and L. Wai-Hsun (Bull. Soc. Chem. 1936, [v], 3, 1389) showed that the following relation applied to a number of organic substances, K=0015 MT, where K was the cryoscopic constant, T the absolute melting-point, and M the molecular weight.

Two main uses of the cryoscopic method must be clearly differentiated. The first as a means of finding approximate molecular weights, which has its main application in organic chemistry, and the second as a method of investigation in the study of the properties of solutions. For the latter work the highest accuracy is proportionality between depression and con- necessary and consequently complicated appara-

an accuracy of a few per cent. is sufficient and test tube, 3-1 cm. in dismeter. The lid and the ease and speed of working are important factors. The Beckmann and Rast methods, which are

of modern precision methods.

THE METHOD OF BECKMANN.—Essentially the procedure is to compare the freezing point of the pure solvent and solution using a thermometer that is calibrated to read differences and not absolute temperatures. Such differences can be measured with much greater accuracy than the actual freezing points, and this fact is utilised in all cryoscopic determinations. The thermometer has a very open scale which covers only 5-6°C, in a length of about 20 cm.; consequently it can be divided into divisions of 1/100°. These divisions can be subdivided by eye to I/I,000° with the aid of a special lens which

elides on the stem of the thermometer and enables errors from parallax to be

avoided. to the upper end of the the quantity of mercury in the hulb may be varied. This permits the for different temperature ranges and hence with various solvents. The adjustment of the amount of mercury in the bulb is carried out as follows. The thermometer is inverted so that the bulb is pointing vertically upwards and gently tapped so that the merand into the reservoir, where it is united with any metal that already

happens to be there. The thermometer is then turned very carefully and the bulb placed in a bath | that a small glass bead wetted with the solvent 2-3°C, shove the highest temperature at which and then cooled in solid carbon dioxide be used it is proposed to work. Provided this operation is done without vibration the mercury will remain joined through the thread from the reservoir to the hulh and auflicient mercury will proviously cooled in liquid air. The freezing flow into the hulh to fill it at the hath tempera-The thermometer is then given a sharp tap with the finger which detaches the excess mercury in the reservoir and the thread will be found to be at the top of the scale at the required temperature. If during the above operation the mercury thread shows a tendency to break, this may be prevented, by holding the thermometer at an angle of 30° to the horizontal, instead of mined as before. One addition, it is true, vertically, while the mercury is flowing from the yields sufficient data for the calculation of the reservoir into the bulb.

A stout glass jar, holding about 11 litres, is on the value obtained for its molecular weight provided with a metal top at the side of which is It is advisable, further, to redetermine the a hole for a stirrer, and in the middle of which freezing-point of the solvent at the end of such a

tus is employed, whereas for the former purpose is another hole capable of holding a large, atout tube are held firmly in position by the springs By means of a cork, there is fitted in the tube capable of an accuracy of about 2-5%, will be another, narrower tabe, to hold the liquid under first described and afterwards an outline given investigation. This tube should preferably have a flat bottom, while the part of it which projects beyond the outer jacket is provided with an auxiliary eide tube, and is itself widened out at the upper extremity. The cork with which this inner tube-the freezing tube, as it may be called-ie fitted carries the thermometer, and allows also free passage to a stirrer. The latter is best made of thin glass rod or of nickel wire. The thermometer must be fitted in an accurately central position, so that the attreer may be moved up and down without touching the hulb.

At the start of an experiment the freezing tube is charged with a weighed quantity of the solvent, usually 10-20 g, and the jer is filled with water or a mixture of ice and salt, or other suitable liquid, at a temperature 4-5° below the freezing point of the solvent under examina-A reservoir is attached tion. The freezing tube, with thermometer and the upper end of the atirrer, is immersed directly in the jar, until the thermometer tube so that temperature has fallen almost to the freezing point. It is then set in the jacket tube and the contents are attrred regularly, while the mercury in the thermometer falls slowly. The fall conthermometer to be used tinues until the solvent is more or less supercooled. In the majority of cases freezing sets in spontaneously before the supercooling exceeds 05°, and the liberation of heat results in a rue of the mercury in the thermometer up to a ateady maximum position. This is the freezing point of the solvent-the temperature at which liquid and solid solvent are in equilibrium with each other. It is advisable, when the thread is rising towards its steady position, to tap the thermometer before taking a reading and so avoid the possibility of the mercury sticking in cury in the bulb begins the capillary. Should the supercooling exceed to flow down the thread 0 5°, crystallisation ought to be initiated. This may sometimes be done by vigorous stirring or, better, hy introducing a small crystal of the solid solvent. D. E. Kenyon and G. A. Hulett (J. Amer. Chem. Soc. 1934, 56, 1649) recommend in place of the crystal, which is rather difficult to introduce. A third method is to touch the surface of the liquid with a thin wire which has been point of the solvent having been determined two or three times in the manner just described, a weighed quantity of the solute, enough to give a depression of not less than 0 2°, is introduced, and the freezing-point of the solution is then ascertained in the same way as that of the solvent. Subsequently additions of solute are made and the resulting depressions determolecular weight of the solute, but it is desirable The other essential parts of Beckmann's to ascertain also whether the concentration of apparatus are seen in the diagram abown above. the solute has any definite and notable influence series of experiments, and so obtain a guarantee that the zero of the thermometer has not changed appreciably in the course of the work.

In the past suggestions have been advanced for improving the accuracy of the Beckmann apparatus. Many of them are now redundant owing to the other types of apparatus developed for accurate work. Consequently only those errors will be considered which affect the apparatus when used for its legitimate purpose-the measurement of molecular weights with reasonable accuracy. The bulb of the thermometer may change slightly in volume during a series of experiments if it has previously been at a considerably different temperature. If this is due to an ageing process in the glass it is best to keep the thermometer temperature as steady as possible. When the bulb is thin pressure changes will affect the position of the thread appreciably. Barometric changes have been shown to have an effect of 0.0002°. The general precautions to be adopted for a Beckmann thermometer to be read with an accuracy of 0.0005° are discussed at length by H. C. S. Snethlage (Rec. trav. chim. 1933, 52, 139).

The temperature of the cooling bath is a matter which has to be carefully considered if the freezing-point depression for a dilute solution is to be correctly determined. Suppose that in an experiment with the ordinary Beckmann apparatus the aqueous solution in the inner tube cools without freezing, it is obvious that the lowest temperature to which the solution can fall must be higher than that of the cooling bath, because of the heat conducted from the air to the solution through the thermometer, stirrer, and tube, and because of the heat generated by the This lowest temperature which the contents of the freezing tube tend to assume when no freezing takes place is known as the convergence temperature. The difference between the convergence temperature and the temperature of the cooling bath varies, of course, with each apparatus; it decreases as the facility for heat exchange between the solution and the external air is lessened, and as the rate of stirring is diminished. Suppose now that the solution under examination has become supercooled, and that it has reached the convergence temperature. The introduction of a crystal of the solid solvent at this stage initiates the freezing process; heat is liberated and the temperature of the solution rises. Inasmuch, however, as the solution is no longer at the convergence temperature, there is an influence at work tending to lower the temperature of the solution, so that the latter never reaches its true freezingpoint; the maximum position reached by the mercury is only what may be called the apparent freezing-point of the solution. Similarly, when the freezing-point of the solution lies below the convergence temperature, an apparent freezing-point is observed which is somewhat higher than the true freezing-point. In both cases the apparent freezing-point lies between the true freezing point and the convergence temperature. In view of these considerations, it is necessary in cryoscopic work to keep the bath

corrected if a value of K is determined with a substance of known molecular weight under conditions identical with those used for the unknown material.

In choosing a solvent for a molecular weight determination it must be remembered that electrolytes in water will dissociate and give low values, whereas many hydroxylated and other organic compounds may associate in benzene (Bury and Jenkins, J.C.S. 1934, 688). When there is a possibility of these complications determinations must be made in more than one solvent. Some work has been done on cryoscopy in mixed solvents, but the results as yet are more of theoretical than practical significance (see O. Anders, Z. physikal. Chem. 1933, 164, 145; G. Scatchard and S. S. Prentiss, J. Amer. Chem. Soo. 1934, 56, 1486).

For the determination of the freezing-point of liquids which are obtainable only in small quantities—physiological fluids, for example—Beckmann's apparatus, in its ordinary form, is too large. Modifications of this apparatus, suitable for the investigation of blood, urine, etc., have been suggested by various authors (Guye and Bogdan, J. Chim. phys. 1903, 1, 379; Burián and Drucker, Zentr. Physiol. 1910, 23, 772; D. Thomas, Z. ges. exp. Med. 1933, 87, 635). In these instruments only 1-2 c.c., or in the last 0·1 c.c., of liquid is required, and it is claimed that the accuracy is of the same order as that attainable with the ordinary Beckmann apparatus.

THE MICRO RAST METHOD.—The determination of molecular weights by the Rast method (Ber. 1922, 55, [B], 1051, 3727) is likely to entirely replace the use of the freezing-point apparatus of Beckmann in the organic chemistry laboratory. It has the advantage of requiring only a very small amount of the solute and the procedure is simpler and quicker than with the older method.

The remarkably high values for the molecular depression of camphor and some related compounds will be noted in the table on page 441d. These substances have excellent solvent action for the majority of organic substances and the fact that there is now a number of them available enables the molecular weight to be checked in two or more solvents, thus guarding against errors due to the separation of mixed crystals of solute and solvent and to association of the solute. Essentially the method consists in observing the melting-points in the ordinary capillary tubes of the pure solvent and of a mixture of known composition. This determination can be carried out in the normal fashion as the depression is large enough for an ordinary thermometer to be used for the temperature measurement. It is best, however, to use a good lens for the examination of the mixture while it is melting. One of two methods of procedure may be adopted, the first when an amount of the order of 0.2 g. of the substance under investigation is available and, second, the true micro-method, which requires only a few milligrams.

is necessary in cryoscopic work to keep the bath temperature as near the freezing-point as stance is accurately weighed into a small test possible. The error is also automatically tube using an ordinary analytical balance.

About ten times this amount of camphor is then highest accuracy at considerable dilution for added, the tube reweighed, and then sealed their confirmation. The precise determinations added, the tube reweighed, and then sealed their confirmation. The precise determination off at the blowpipe. The tube is heated in a for recent years have all been done in development as the emperature alightly above that of the members of a type of apparatus evolved hy adome until all the solute has dissolved. The tube is two Deway vessels, one of which contained removed from the bath and shaken while it so Deway vessels, one of which contained contents powdered in a small agate mortal and the determination. The mounts of the years were closed with corks Samples of this powder are placed in ordinary and they were supported in a metal can which was nature point capillary tubes and the determination. perature being noted at which the last trace of solid disappears. The melting-point of the water or ice and solution required that there be compher alone is then determined and from the efficient circulation of the liquid. This was Compion Alone is included a separation of the solute conditions of the condition of the con for the sampla used. Benzoic acid or naphtha equilibrium was rapidly established. The lens are the materials that are commonly used temperature difference between the two Dewar for standardisation.

method some knowledge of micro-chemical manipulations and the use of a micro halance are required. A capillary tube 40 mm. long and 2-3 mm. in diameter is weighed to 0 000001 g. shout 02 mg. of the unknown substance is placed in the closed end of the tube which is then reweighed, ten times the amount of camphor is introduced and its weight also found. The capillary is then drawn off at about 15-20 mm from the closed end, the glass being formed into a long fibre by which the small hulh containing the charge can be held. The camphor and the solute are melted together until the latter has completely descrived and the melting-point of the mixture is then determined. For details of these operations F. Pregl's " Quantitative Micro analysis" may be consulted (3rd ed., 1937, J. & A. Churchill, Ltd.).

J. Purseh has been responsible for many improvements in the Rast method in recent years. He has described a modified procedure to be adopted when the molecular weights of volatile compounds are required (Ber. 1932, 65, [B], 1227); also a large number of naw solvents have been examined. These are recommended to their the results obtained with tampion and to be used when the substance is insoluble in, or decomposes at, the melting point of camphor, The more important of these materials are:

Camphene, m p. 49°C., K=310 (Pirsch, Ber. 1932, 65, [B], 862).

Bornylamine, m.p. 190°C., K=406 (Pirsch, Ber. 1932, 65, [B], 1227). (Specially useful for alkaloids and basic substances.

Camphorquinona m p. 190°C, K=457 (Ber. 1933, 66, [B], 815).

cycloPentadecanone, m p. 65 6°C., K=213 (Giral, Anal. Fis. Quim. 1935, 83, 438). (Used for azo dyes, also for sterols and derivatives.)

been used will be found in the following papers 1925, 58, [B], 2388; Bul. Chim. 1926, [4], 39, 82: Rocz. Chem. 1933, 13, 7201,

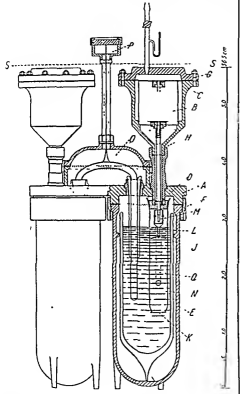
with ice. The equilibrium between ice and water or ice and solution required that there be vessels was measured by a 50-junction thermo-For the successful operation of the second couple of copper constantan, the wire of which ethod some knowledge of micro-chemical was tested for homogeneity. When the E.M.F. was read to one microvolt the temperature could be determined to 0 0001°. After equilibrium had been established some of the solution was withdrawn and its concentration determmed with the aid of an interferometer, solutions of nearly equal and known concentration being used for comparison. In this way determinations were carried out with solutions as dilute as 0 004 normal.

F. Hovorka and W. H. Rodehush (J. Amer. Chem. Soc. 1925, 47, 1614) introduced the following refinements into their apparatus. The mechanical stirrer was replaced by a stream of purified air which was found to give efficient mixing and not cause so much heating as the previous methods. Also, instead of withdrawing and analysing a sample of the hourd, the solution was made up to a known concentration and cooled with the rest of the apparatus to a temperature very close to that of the equilibrium The solution was then added to the ice in the Dewar vessel from which the water had been withdrawn a few moments earlier. In this way the equilibrium temperature was reached without upprevaile thange in the tencentrium of the solution. The temperature difference was measured with a 14-junction thermoelement, and improvements were introduced into the circuit for measuring the potential and excluding parasitic E.M.F.'s. This rendered it possible to read the temperature with a precision of 0 00002°.

Randall and Scott (J. Amer. Chem. Soc. 1927, 49, 647; Randall and Vanselow, shid. 1924, 46, 2418) surrounded the Dewar flasks with aubetantial metal casing which enabled the whole apparatus to be exhausted to the vapour pressure of the water, thus excluding the error that may Descriptions of other materials that have be introduced by changes in the solubility of the en used will be found in the following papers air with temperature. The concentration of the (Ber. 1933, 66, [B], 1694; 1934, 67, [B], 1115; solution was found from conductivity measure. menta. L. Ehert and J. Langa (Z. physikal Chem. 1930, 149, 389; 1934, 168, 147) lisve PRECISION CRYOSCOPY. - The modern theories introduced further improvements, their final of solution require experimental results of the apparatus being shown in the figure on p. 445.

device M for preventing undue conduction of heat by means of the induction motor B. Since the armature of the motor is enclosed in the casing there is no need for packed joints with consequent leakage. The thermocouple O is enclosed for the greater part of its length in a Dewar vessel which minimises the conduction of heat down the elements. The concentration of the solution is again found by a conductivity method. Using a 30-junction thermo-element the temperature could be determined to 0.00001°.

Scatchard and Prentiss have also made a series of careful measurements for a variety of alkali



salt solutions. Their apparatus is not evacuated but filled with nitrogen which they say does not introduce appreciable errors (see J. Amer. Chem. Soc. 1932, 54, 2676, 2690, 2696; 1933, 55, 4355; 1934, 56, 1486, 2314, 2320; 1936, 58, 837).

The lowering of the freezing-point of nonaqueous solutions has also been considerably developed. L. D. Elliott (J. Physical Chem. 1924, 28, 611) described an apparatus in which liquid ammonia can be used. The design of the apparatus is similar to that of the ordinary Beckmann apparatus except that the cooling bath is contained in a large Dewar vessel and that arrangements are made for cooling the light petroleum it contains with liquid air. At these low temperatures a resistance thermometer is the best instrument for measuring the temperature. Measurements for ammonium nitrate and sulphurio acid in anhydrous acetic characteristic angles.

A mechanical stirrer J is driven through a acid have been carried out with an accuracy of 0.0001°C. by W. C. Eichelberger (J. Amer. Chem. Soc. 1934, 56, 799). F. S. Brown and C. R. Bury (J. Physical Chem. 1926, 30, 694) have studied the depression of the freezing-point of nitrobenzene by organic compounds. For work with cyclohexanol, see E. Schreiner, O. E. Frivold and F. Ender (Phil. Mag. 1929, 8, 669); with indene, see W. Klatt (Z. physikal. Chem. 1934, 171, 454); and with ether-alcohol mixtures, see A. Lalande (Compt. rend. 1932, 195, 1267). F. Garelli and E. Monath (Atti R. Accad. Sci. Torino, 1926, 61, 4) have suggested that the second series of the seco that the cryoscopic method might be employed for the determination of the solubility of gases in organic liquids, and they have determined the solubility of nitric oxide and carbon monoxide in benzene, bromoform and nitrobenzene by this method.

> R. H. P. CRYPTAL. An aldehyde, to which the name cryptal was given, was stated to occur in the oils from Eucalyptus hemiphloia, polybractea and Bakeri (Penfold, J.C.S. 1922, 121, 266; Penfold and Simonsen, ibid. 1930, 403). This substance has now been shown by Berry, Macbeth and Swanson (ibid. 1937, 986) to be identical with the ketone, l-1-4-isopropyl-\(\Delta^2\)-cyclohexen-1-one isolated from these oils by Cahn, Penfold and Simonsen (ibid. 1931, 1366).

> J. L. S. CRYPTOLITE v. CERIUM METALS AND EARTHS, Monazite.

> CRYPTOPINE, an isoquinoline alkaloid. CRYPTOTAENENE. \mathbf{The} hydrocarbon, cryptotaenene, $C_{10}H_{16}$, b.p. 67-68°/15 mm., d_4^{25} 0·8128, n_D^{25} 1·47476, $[a]_D$ +2·66° occurs in the essential oil obtained from Crypto. taenia japonica (Hirano, J. Soc. Chem. Ind. Japan, 1926, 29, 48). It contains three ethylenic linkages and gives on oxidation with ozone. acetone, formic and succinic acids. stitution has not been determined.

J. L. S. CRYPTOXANTHIN 17. CAROTENOIDS.

Kryptoxanthin (Vol. II, p. 399c). CRYSTALLISATION. The researches of William L. Bragg ("Crystallisation," Trans. Inst. Chem. Eng., 1927, 5, 103-109, and other publications) and others into the crystalline structure of matter have shown that in a crystal the molecules of which it is composed are arranged in a definite space lattice structure and are capable only of oscillatory movement, so that crystalline structure possesses great rigidity and large stresses must be applied to produce plastic flow.

The spacing of the molecules in the lattice, although regular, is not the same in all directions and therefore the forces of attraction between the molecules will vary and result in the crystal possessing different properties along various

Each crystalline substance has its own characteristic arrangement of the space lattice structure and if the crystal is allowed to form freely it possesses a definite geometrical form having a fixed number of plane surfaces which are inclined to one another at definite and

Many substances are known to adopt two or to the attraction of the molecules in the more wholly different structures depending upon the physical conditions which are prevalent force becomes the greater and the molecule during the growth of the crystal and are termed tends to enter the solution, but if the vinction polymorphic.

Sometimes the conditions required to produce a change in structure are well understood and when they are applied a crystal is obtained which is perfectly stable so long as the conditions, such as a definite transition temperature, are maintained. In other cases the conditions governing the formation of different structures are obscure, the rarer structures heing unstable and the crystal either dissolving in the solution or breaking into minute aggregates of the

commoner type. With organie substances it is often difficult to decide whether the modifications in the crystals of a substance are due to polymorphism or to isomerism, that is, permanent intra molecular

differences in structure.

To distinguish between polymorphism and isomerism Sidgwick's (J.C.S. 1915, 107, 672) method of studying the equilibrium relations between the two forms and a common solvent can be applied, when it will be found that with isomerides differences will survivo a process of

either solution or fusion. Crystal Growth -If it is assumed that the forces of attraction between the molecules constituting the crystal act along hies joining tha centres of the molecules, then a diagram can be constructed showing the vectorial distribution of the surrounding field Molecules within tha crystal will be subjected to a balanced system of forces due to the attractions of the surrounding molecules, whereas a molecule at the surface of the crystal will be acted upon by an unbalanced force which tends to attract other molecules to itself or produce a strain at the free surface of the crystal. This surface energy depends upon the number, kind and distribution of the molecules on the plane surface and may vary for different surfaces of the crystal Moreover, molecules at the corners and edges of the crystal are subjected to smaller attractive forces towards the centre of the crystal and so cannot adhere so strongly to the crystal as those at or near the interior of the plane surfaces. J. Willard Gibbs ("Scientific Papers," London, 1906, p. 325) has indicated that these factors supply much information on the mechanism of crystal growth When a crystal is immersed in its saturated solution and a state of equilibrium between the crystal and solution is attained the number of molecules. arriving at the surfaces of the crystal from the solution must equal the number of molecules leaving these surfaces and entering the surrounding liquid.

A molecule vibrating about an equilibrium position along a line at right angles to the plane surface of the crystal will be subject to an attractive force towards the crystal due to the other molecules forming the crystal and an outward force of attraction due to the molecules constituting the surrounding liquid. If the form spinteneously in superstaturated solutions, with my reach a point termed the halance point are the halance point. Trans. 1903, 202, 459-623, Proc. Roy. Soc. where the attraction due to the cryetal is equal.

of the molecule is not sufficient to carry it to the balance point then the attraction towards the centre of the crystal heing the greater force. the molecule will tend to return to the curface of the crystal.

The same theory also applies to the molecules of the solute in the surrounding liquid which not being aubject to a directional movement may or may not come sufficiently close to the curface of the crystal to be attracted towards it. greater the number of the molecules of solute present in the solution surrounding the crystal the greater will be the tendency for these to be attracted to the crystal surfaces. Hence the concentration of the solution is another impor-

tant factor in crystal growth,

Other factors, in addition to the kinetic energy of the molecule, such as inter-molecular collisions which are dependent upon the number of molecules in the surface layer, and the periodicity of the vibration as well as the designation of the heat of solution and the change in density of the solution affect the growing or dissolving of a crystal. As the attractive forces on the molecules at the edgee and cornere of the crystal towards the centre of the crystal are least the amplitude of their vibrations will be greater than those at the interior of the plane surfaces and therefore the tendency will be for those molecules to leave the crystal and enter the solution Hence, during the growth of a crystal, complete molecular planes must be formed on the surfaces, and since single molecules at the corners edges or on the surface are held less firmly a definite degree of supersaturation of the surrounding solution is necessary to mainten erystal growth. Before, however, crystal growth can occur there must be a solid nucleus upon which the molecules can be deposited.

Nucleation -The nucleue upon which the molecules are deposited to form the crystal may consist of a small solid particle inserted in the hourd or may arise by formation of euch a particle in the solution by an agglomeration of a number of molecules of the same substance of which the creatal will be composed, to give a small solid particle upon which the molecules can be

deposited.

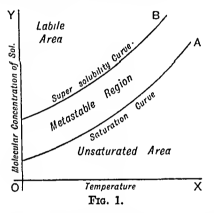
The introduction of small crystals of the cubstance to be crystallised into the solution in which the gicwith takes place is known as the seeding of the solution and is cometimes adopted in practice, but the commoner procedure is to form the solid nucleus from the molecules in the solution,

Since the attraction towards the centre of the crystal is least for molecules at the corners or edges, if the particle constituting the nucleus consists of only a few molecules the tendency for these to reenter the solution ie greatly mereased (G A. Hulett, Z. physikal Chem. 1901, 37, 385), and therefore such nuclei can only

1908, 93, 825) by their researches on the concentration of solutions during the progress of crystallisation have contributed materially to our knowledge of the conditions under which nuclei will form spontaneously in a solution, and the requirements which must be fulfilled if

crystals of uniform size are desired.

From his observations Miers found that in addition to the ordinary solubility curve A, which divides the area between the X and Y axes into two portions such that if, at any given temperature, the molecular concentration of the solution is represented by a point below the curve A the solution will be unsaturated, if on the curve A saturated, and if above the curve A supersaturated, a second curve B, which has been termed the supersolubility curve, could be drawn which divides the supersaturated area of the diagram into two portions.



The region between the saturation curve A and the supersolubility curve B is one of metastable equilibrium, whilst the region above the supersolubility curve is one of unstable-equilibrium.

If the temperature and molecular concentration of a solution be such that it will be represented by a point in the metastable then it will be found that nucleation will not occur, but if such a solution is "seeded" then these particles will tend to increase in size but not in number.

The metastable region therefore is favourable to the growth of a crystal but not to the formation of the nucleus upon which the growth takes place. On the other hand, if the temperature and molecular concentration of a solution are such as to be represented by a point above the super-solubility curve in the labile region, then it will be found that as long as these conditions are maintained nucleation and crystal growth will occur simultaneously. will result in a product consisting of a large number of crystals of varying sizes, since the nuclei first formed, being for a longer time in contact with the supersaturated solution, should have grown to a larger size than subsequent nuclei; the smallest crystals being obtained from the nuclei last formed.

The formation of nuclei by the cooling of a pure solution appears to be a characteristic quality of the molecules, for in some cases only

slight undercooling will cause nucleation, whereas with other solutions great undercooling must be employed before the nucleus is formed. As the degree of undercooling increases, nucleation occurs more readily until a maximum is reached, but after this point intense undercooling results in diminished nucleation. Since the mean kinetic energy of the molecules diminishes with fall in temperature if this were the only factor, rate of nucleation would increase with lowering of temperature. Viscosity, on the other hand, regularly increases as the temperature is lowered, and it is therefore assumed that when maximum nucleation takes place the kinetic energy of the molecules and the viscosity of the solution are particularly favourable to the change from isoto anisotropic grouping. At lower temperatures the greater vicsosity of the liquid inhibits the change which is also hindered at temperatures above the maximum point by the larger movements of the molecules.

With very rapid cooling it is possible to pass quickly through this point of maximum nucleation and reach a region where the viscosity is so great that crystal formation becomes impossible and when the solution will possess both high viscosity, great rigidity and, at the

same time, be isotropic in character.

The rate of nucleation is also greatly modified by the presence of even small quantities of impurities in the solution, some impurities even arresting the formation of the nuclei and others greatly increasing the degree of nucleation (G. Tammann, "The States of Aggregation," New York, 1925, pp. 225–251).

Rapid cooling of the solution usually results

Rapid cooling of the solution usually results in the production of a very large number of very small crystals and industrially this is sometimes of great advantage as in the preservation of food, where in the cold storage of food large crystals would tend to produce rupture of the cell walls.

Crystallisation therefore depends upon two principal factors: (a) nucleus formation, and (b) subsequent growth of the crystal on the nucleus, and is employed on an industrial scale:

(1) to remove a dissolved substance from solution;

(2) to separate a dissolved substance from a solution containing other substances;

(3) to produce a dissolved substance from a solution in a marketable form.

(1) Removal.—The removal of a dissolved substance from solution can in many cases be effected more economically by evaporation to dryness and where crystallising plant has been installed to remove a dissolved substance its installation was attributable to equipment suitable for complete evaporation not being obtainable.

There are, however, instances in which evaporation to dryness, even with modern equipment, is an uneconomical or impossible means of recovering the dissolved material because of a factor such as the operating temperature being too high, in such cases crystallisation has to be used.

(2) Separation.—When the object of the crystallisation process is to separate one

be given regarding procedure, for a factor of where they are supported by the surface tension great importance is the solubility relationships of the liquid until they have grown so large

(3) Production of Marketable Crystals.—Here the process of crystallisation is employed to suspended those surfaces of the crystal which are produce from a solution, in which it has been completely submerged in the liquid will grow in dissolved, a solid material for which there is a

ready sale.

Within recent years crystals of nniform size have commanded a better market than consignments of the same material in which the sizes of the individual crystals vary greatly. A reason for this demand is that adulteration of the cooling is effected by (a) evaporation at the a quantity of crystals of uniform size is more easily detected then of a similar consignment through the walls of the vessel, and the tem. consisting of crystals of widely varying sizes, because in the former ease the adulterant, if it is not to be easily detected by the eye, must consist of crystals having the same colour, shape and size as the other crystals.

Crystallising Plants -The production crystals of a material from a solution in which it is dissolved can be effected in several ways by, (1) cooling the hot saturated solution ;

(2) evaporating a solution until the concentration of the dissolved substance is such that it will either promote nucleation and maintain subsequent growth of the nucles or maintain the growth of seeding crystals added to the concentrated solution. This may be carried out at varying pressures either above or below atmospheric. Sometimes evaporation and cooling are combined in order to achieve the desired result: or

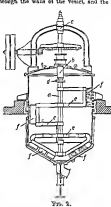
(3) other methods, such as by addition of materials which after the solubility of the dusolved substance in the solution, etc.

CRYSTALLISATION BY COOLING.

The crystallising pan, one of the earliest forms of crystallisers in which uncontrolled crystallisa. tion by cooling is carried out, is a long rectangular trough made of materials which will resist the action of the solution from which the erystals have to be formed or if attacked by this solution will not spoil the colour of the crystals obtained or man the product by the presence of too much impurity Such tanks are often made of cast iron or of wood haed with lead. tm, zinc or other austable metal, and provided with rounded corners, and having the bottom of the tank sloping towards the nutlet. Into such a vessel is run the mother liquor at such a temperature and concentration, which have been previously determined for the particular solution, as will yield the best results. When cooling is complete the liquid is run off and the crystals which are usually adhering to the sides and bottom of the tank removed by which is fixed the outlet valve and pipe. hand or mechanical acrapers. Sometimes threads or wires are suspended from rods placed a shaft a driven by bevel gearing mounted on across the tanks upon which crystals are formed, the cover of the crystaliser which is also although when threads are used their entangles, provided with an inlet for the mother liquor, ment, produced by convection currents in the cooling liquid, often results in large crystals of it the solvent is a valuable or harmful material. very arregular shapes. Often, too, with this an opening can be arranged for the removal and method of crystallisation, irregular crystals are recovery of the vapour. Since the crystals

constituent from a solution containing two or produced by the convection currents carrying more dissolved substances no general rules can very small crystals to the surface of the bond very small crystals to the surface of the liquid of the various substances present m the solution. that their weight is greater than this supporting force. During the time that they are thus a normal manner, whilst the growth on the other surfaces which are not completely in contact with the liquor will be abnormal. The shape, therefore, of such a crystal will not be regular.

Since in the crystallising pan described above free surface of the Liquid or (b) by conduction



perature gradient across the walls is chiefly maintained by the natural convection currents set up within the liquid as it cools, the cooling in this manner is very slow. To increase the loss of heat from the sides of the vessel and so merease the rate of cooling mechanical agitation is often employed. Such a plant for uncontrolled crystallisation by cooling shown in Fig 2, consists of a cylindrical vessel with an inverted conical base at the lowest point of

Within the cylinder is an agitator's carried by

usually form on the inner surface of the cylindrical shell and reduce the flow of heat from the vessel, Fig. 2 also illustrates a method, namely, by attaching brushes or scrapers f to the agitator, of preventing this increased resistance to the flow of heat through the cylindrical walls. The lower portion of this crystalliser is also provided with a jacket through which a cooling liquid can be circulated, and if a highly concentrated solution has to be cooled a strong agitator must be fitted, and in some instances means provided for the removal of the crystals shortly after they are formed.

Another crystallising apparatus, the Passburg, suitable for producing low temperatures such as are used in the formation of crystals of potassium chlorate, is shown in Fig. 3. In this plant the mother liquor enters the left hand

the cooling liquid circulates and to maintain a high coefficient of heat transmission is fitted internally with a set of revolving brushes, J, which remove the crystals from the cooled walls. They then fall into the conical base of this cooler. from which they can be discharged through the valve L into the centrifugal. The cooled mother liquor leaves near the bottom of the vessel, and passing upwards through the inclined pipe M is discharged into the right-hand cooler D. Into this cylinder is fitted a spiral coil C, the lower end of which is coupled to the cooler B, which is connected with the pressure side of the compressor A. The outlet of this coil is at the top and is connected by the pipe E with the suction valves of the compressor; the coil, cooler, and compressor thus constituting a refrigerating system. Brushes arranged on a central shaft crystallising vessel through the pipe K. This which is rotated through suitable gearing P vessel is provided with a jacket through which fixed to the cover of this vessel remove the

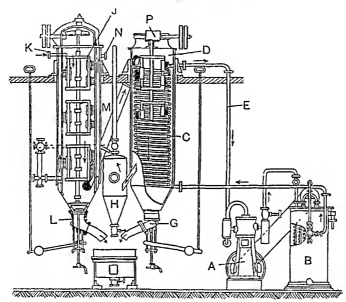


Fig. 3.—Passburg Crystalliser.

crystals from the surface of the cooling coil. A conical cover is fitted at the bottom of this cooler which collects the crystals removed from the cooling surface. By opening the valve G at the lowest point of this cover, these crystals, together with some of the mother liquor, can enter the centrifugal where separation is effected. The cold liquid leaves this cooler at the bottom and passes to the bottom of a smaller cylinder, H, also fitted with a conical base. having a valve at the lowest point. Whilst slowly ascending this cylinder some crystal growth occurs and this crop of crystals together with some mother liquor is, at regular intervals, discharged into the centrifugal. On reaching the top of this cylinder the mother liquor passes through a pipe into the foot of the jacket of the first cooler. After ascending this jacket and thereby cooling the incoming liquid this cooling solution leaves the crystalliser by the pipe N.

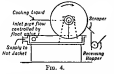
intermittently and are sometimes designated batch erystallisers, but there are others, which, still adopting the principle of cooling the

solution, operate continuously.

Sometimes crystallisers belonging to this group consist of a trough having a semicylindrical base and fitted with a spiral agitator mounted on and driven by a shaft the axis of which coincides with the axis of the trough. In another type of uncontrolled crystalliser the hot concentrated solution is run into a shallow, rectangular tray. Dipping into this tray is a hollow cylinder with its axis horizontal. Through the trunnions supporting this cylinder pass pipes for the admission and removal of the cooling liquid. By means of a ball-valve the hot concentrated solution in the tray is kept at such a level that the rotation of the drum causes its outer surface to be covered by a thin film of liquid from the tray. Contact with this cooled The crystallisers described above operate surface causes the crystals to form on the surface

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of the drum from which they are removed by | which extends tha whole length of the trough acrapers or doctors placed as shown in Fig. 4, and carries the worm or ribbon conveyor C. which gives an end view of the film or dipping being driven by means of a worm and worm drum type of crystalliser which is continuous wheel D supported on the end at which the hot



Amonter Communicative and the second before the solution and crystals formed are Walker crystaliser, Fig. 5, which consists of an finally separated usually by centrifuging A open trough Javing a semi cylindrical base variant of this type of crystalisers cometimes

in operation. The examples given illustrata solution is admitted. Surrounding the lower portion of the trough is a jacket I through which the cooling liquid can circulate. These crystallisers are usually made in sections 10 ft. long, four sections being boited together to produce a trough 40 ft. in length. The outlet F of the jacket of one section is connected by an inverted U tube, G, to the inlet of the next section, H, the process being repeated until all the jackets are connected in series. Countercurrent flow of the hot solution and the cooling hand is thus assured. The end plate of the last section is fitted with a weir, K, over which the cooled liquid and crystals flow, either into a Another continuous crystalliser employing similar crystallisers for meries through one or more similar crystallisers if further cooling is desired by the Swenson before the solution and crystalliser. Fig. 5. which constitutes the cooling is desired the solution and crystallisers. eentrifugal machina if crystallisation has been and placed with its axis horizontal. The axis of adopted is a long tube which is mounted and the trough coincides with the axis of a shaft B, driven in a similar manner to a tube mill. The

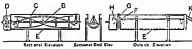


Fig. 5 .- First and Last Sections of Swenson-Walker Crystalliser.

nerves as two vature, a stationary errow | use inghest point, in its passage through the conveyor or plates arranged spirally on the crystalliser must follow a simuous path before it may be surface of the tube being comployed to reaches the outlet placed in the lower end plate.

Rotating conveyors in these crystallisers unless carefully adjusted are hable to break a number suitably controlled hydraulic rams or by

of the crystals thus forming additional nucles and so give an output of crystals

of varying sizes.

In an attempt to redure this attrition of the crystals to a minimum and so produce a uniform product, the Wulff and Bock or rocking crystalliser has been developed.

Such a machine consists of a long trough up to 100 ft. in length and supported so that its axis is slightly inclined to the horizontal, Fig. 6 represents a view of such a crystalliser and ahows the rockers or rings, tha rollers which support them, and that bracing between the trough and the rockers and each pair of rockers. The trough is designed so that as it is rocked to and fro the solution will not be ejected over the edges and is fitted with a series of transversa vertical

nately on the right and left hand sides of the etructure.

hot solution enters at one end and the cooled trough the solution, which is admitted at solution leaves at the other, a stationary screw the highest point, in its passage through the

Rocking of these crystallisers is effected by

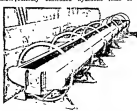


Fig. 6 .- Wulff and Bock Crystalliser.

befile plates which divids the length of the electrically driven rocking gear. Special attentionals into a series of compartments. As these tion abould be given to the attength of the partitions do not extend across the trough machine because at the point of reversil of but are arranged to provide openings eller-

Another crystalliser in which crystallisation is effected by cooling a saturated solution whilst in motion is the Howard Crystalliser, of which Fig. 7 represents a diagrammatic view. This apparatus is formed of three truncated conical vessels placed one above the other and having their axes in line. The lower two vessels are provided with jackets, whilst in the upper vessel there is a coiled tube. The saturated solution is admitted to the lowest vessel B through the pipe A, and after filling this vessel rises into the middle vessel D through the short tube C.

Inside this vessel is another conical chamber, E, within which is a pipe, for the removal of cooling liquid, which extends nearly to the

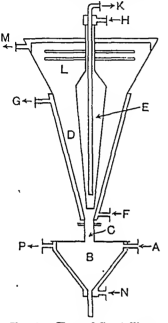


Fig. 7.—Howard Crystalliser.

bottom of this vessel. The taper of this conical vessel is such that in the ascent of vessel D the sectional area of the annulus formed by these two vessels continually increases. In this way the velocity of the liquid rising in vessel D is being continually reduced and so only crystals of a definite size are capable of falling into the vessel B. Cooling liquid is admitted to the jacket of vessel D at F, and removed at G, whilst the cooling liquid for vessel E is admitted at H and removed at K. On reaching the top section, L, crystal growth is stopped by circulating a hot liquid through the coils which heats the surrounding liquid before it flows over the weir fixed at the top of this vessel and so reaches the outlet M. Hot liquid is also circulated through the jacket of vessel B, entering at N and leaving at P, and this maintains the temperature of the liquid in B at such a point that no further growth can take place on the crystals in this vessel. The lower end of B is connected to a salt box or boot of an elevator by which the crystals can be removed from the plant.

Formation of crystals by hot solutions can also be effected by forcing the solution in the form of a fine spray into a cooling tower or by making the solution fall in the form of drops through a current of cold gas or air which is drawn through a duct (" Der Chemie Ingenieur," vol. 1, part III, pp. 203, 204).

CRYSTALLISATION BY EVAPORATION.

An early type of plant for this purpose resembles the tank crystalliser used for uncontrolled crystallisation by cooling, but in this case the tank is arranged in a brickwork setting so that hot gases from a furnace placed in the setting at one end of the tank can pass over the walls and bottom of the tank and heat its contents. Sometimes these tanks are provided with a series of parallel tubes placed near the bottom of the tank and fixed between the end plates of the tank through which the hot gases from the furnace, placed at one end of the tank can pass on their way to the chimney. During both the cooling and evaporating methods employed in crystallising pans, vapour is evolved at the free surface of the liquid and if still air conditions are maintained, above the pan, the following formula, devised by J. W. Hinchley, enables the rate of evaporation of the solution to be determined:

$$W = A \left(\frac{p_e - p_d}{50}\right)^{1 \cdot 2}$$

where W=weight of liquid evaporated in kilograms per hour per sq. metre of surface of the liquid; p_e =vapour pressure of the solution in mm. of mercury at the temperature under consideration; p_d =pressure of the aqueous vapour in the atmosphere of the factory; and A=constant, depending on the solution undergoing evaporation.

If air currents are directed over the free surface of the liquid then the work by Hinchley and Himus (Trans. Inst. Chem. Eng. 1924, 2, 57) and other workers (Himus, ibid. 1929, 7, 166; Powell and Griffiths, ibid. 1935, 13, 175) shows how the evaporation can be determined for a given set of conditions. With this type of crystalliser the movement of the liquid is largely due to convection currents, and therefore the growth of small crystals, which are retained at the free surface by the surface tension of the liquid, will not occur in a regular manner.

Evaporators can be used to concentrate the solution, which may then pass into a cooler, or a vacuum pan, where the crystallisation is carried out, or the arrangement may be such that the crystals are formed in the evaporator or separator of the plant and are subsequently withdrawn after the removal of the surplus mother liquor. Usually the latter type of plant is referred to as a salting evaporator and has a special field in the production of fine-grained crystals.

When evaporators are used to concentrate the solution either for the coolers or vacuum pans any of the different types of heating may be employed, but in the salting evaporators vertical tubes are used in the heating element. The commonest forms, are a series of vertical tubes

central downcomer, the basket type of heater, plato upon which a quantity of crystals have been and the individual heater or calandria of the climbing film type. In the first two types the the liquid becomes superheated due to increased lower cover attached to the evaporator shell is conical and its lowest point connected with the device for removing the crystals. Sometimes when a central downcomer is employed forced circulation of the liquor in the evaporator is produced by means of a propeller placed at er near the bottom of this tube In the consent base below the hesters is a region in which the solution is comparatively quiescent and in which growth of erystals can proceed until they are able to fall into the receiver When a separate calandria or heater is employed the hot liquor has to be discharged into a separator where the vapour produced during its passage

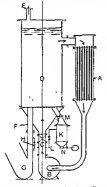


Fig. 8.—Borake Saltverker Crystalliser,

through the heater or generated from the super heated hour by the reduction in pressure, is separated from the hour. At the bottem of this separator means are provided for the recovery and removal of the crystals produced and the return of the liquid to the calandria for further heating.

An important development in the subject of crystallisation by evaporation was published in Chem Trade J. and C. Eng. Jan. 16, 1925, when a crystalliser developed by the A. S. de Borske Saltverker of Bergen for the production passage through the heater A is forced by the of coarse grained salt for fishery work was centrifugal pump B into the vessel C, where described.

Fig. 8 illustrates diagrammatically the early reduction in pressure. form of this crystalliser. In this arrangement of the plant the solution is drawn through the at this point results in the solution becoming heater A by the pump B and forced through the supersaturated and this fills by the pupe D valve C into the bottom of the chamber D. into the crystalliser E. As the lower end

extending between two tube plates with a large At the bottom of this chamber is a perforated placed. During its passage through the hester pressure produced by the increase in the head of the liquid and no vapour is generated until it is near to the free surface of the bould in D. If the solution is saturated the evolution of vapour, which escapes through E, causes the remainder of the liquid to become supersaturated. Part of this again passes through the heater and is again superheated, the cycle being thus continuous. Thus the crystals on the perforated plate are always in contact with a supersaturated solution and will increase in size Moreover, the bound from the centrifugal moving upwards will keep these in suspension until they attam such dimensions that they can overcome the npthrust of the rising liquid, when they will fall into the tubo F connected with the shoe of the hucket elevator G, by which they are removed from the crystalliser. Liquid is fed continuously to the crystalliser through the pipe H, and hy adjusting the valves on the pipe H and the discharge from the centrifugal pump C. the upward flow of liquid in the pipe F and through the perferated plate can be regulated so that only crystals of a definite size fall into the shoe of the elevator,

An alternative method of removing tha erystals, employing crystal hoves, is shown at K. If desired several of these can be arranged so as to remove the crystal from soveral points round the perforated plate. Here again the feed which is admitted by the pipe L provides the upward flow which regulates the size of crystal entering the hox. When several boxes are eroployed, then by closing the feed valve and the valve M and opening the valve N the crystals can be discharged from each box in turn and thus produce the minimum disturbance

in the running of the plant The hucket elevator and the crystal box are two common methods of removing the crystals from salting evaporators. When the hucket elevator is used the casing enclosing the buckets must be liquid tight and must extend to such a height that the hydrostatic pressure produced by the head of cool liquid above the hottom of the choe will always be equal to the pressure on the free surface of the liquid plus the pressure due to the height of the hot liquid measured from the same datum level Two or more crystal boxes are usually provided, so that whilst one is being emptied of its crystals the crystals which are being formed can collect in the others.

A modified form of the crystalliser shown in Fig. 8 is the Jeromiassen or Oslo crystalliser (J II. Perry, "Chemical Engineering Hand book," 1934, 1884), Fig. 9. In this plant the liquid, after being superheated in its downward evaporation takes place as the result of the

Removal of solvent from a saturated solution

of the pipe D passes through a perforated plate fixed near the bottom of the erystalliser and upon which the seed crystals are supported, the supersaturated solution discharged by D must by its upward passage through the plate maintain the crystals thereon in suspension until they attain such dimensions that they can

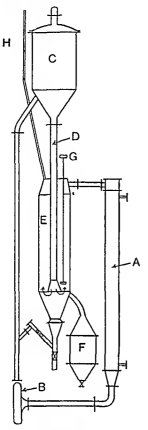


Fig. 9.—Jeremiassen or Oslo Crystalliser.

fall against the ascending stream on to the perforated plate. On opening the valve G these crystals enter the crystal box F, and can be withdrawn from the crystalliser whilst the solution rising to the top of the crystalliser again is superheated by passing through the heater A. Liquid is supplied to the crystalliser through the pipe H, the feed being either of a continuous or intermittent nature.

CRYSTALLISATION UNDER REDUCED Pressure.

In some industries, notably the sugar industry, the concentrated liquid is pumped into a vacuum pan where the final water is evaporated under reduced pressure and the crystal formation

Vacuum pans closely resemble evaporators, the heating elements being either coils or one of the various types of tubular heaters. Syrup

so that it becomes practically impossible for the crystal of sugar to settle in the liquor and therefore special attention must be given to maintaining good circulation of the liquid in the pan.

When eoils are used for heating the liquid the spacing between the top coils should be greatest and gradually diminish in the lower layers. Heaters having large vertical tubes also materially assist circulation, whilst an essential feature of such pans is a downcomer of ample cross-sectional area.

Another important factor in operating such pans is the vacuum, for it is generally conceded that a high vacuum results in erystals of soft grain whilst hard grain crystals are obtained if a low vacuum is employed. Usually in the sugar industry the working pressure in the vapour space of the vacuum pan is below 31 lh. per sq. in. absoluto. If the viscosity of the saturated solution is low, then in crystallisers of this type small crystals are produced because the crystal once formed tends to sink rapidly through the solution. To produce large crystals in such a plant the problem is largely to maintain these small erystals in contact with the supersaturated solution for sufficient time to allow of their

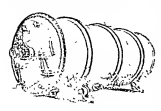
growth to the size required.

In the vacuum crystalliser, however, another principle is involved. If a saturated solution at a temperature corresponding to its boiling-point at atmospheric or higher pressure is admitted to a vessel within which there is a lower pressure. the solution will be at a higher temperature than a similar solution boiling at the lower pressure This superheated liquid will fall in temperature to the boiling-point of the liquid at the lower pressure and the heat thus rendered available will eause some of the solvent to vaporise. The remainder of the solution will therefore become supersaturated and crystallisation or crystal growth will occur. Such a method cools the solution without transferring the heat to be dissipated through the walls of the container, and can be used for cooling nearly all aqueous solutions and many solutions in which organic solvents are employed. Condensation of the vapour evolved in a suitable condenser maintains the requisite pressure in the crystalliser and allows the solvent to be recovered. This is particularly advantageous when organic solvents are employed, especially in large-scale operations. Fig. 10 (Griffiths, Chem. Eng. Group 1924, 6b. 9) illustrates a crystalliser of the rocker type, the hot saturated solution being admitted by a pipe passing through the end plate at the higher end, whilst the cooled solution together with the crystals are removed through suitable openings in tho lower end plate. Connections are also provided on this plato for the removal of the vapour evolved.

Another vacuum crystalliser (Badger and McCabe, "Elements of Chemical Engineering," 1931, p. 418) is shown diagrammatically in Fig. 11. Here the hot saturated solution enters at A into the vessel B in which the pressure is less than atmospheric and as a result an amount of vapour is evolved which is removed through C to the condenser or vacuum pump by which fed to these vacuum pans has a high viscosity, the reduced pressure in B is maintained. The

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supersaturated solution thus formed permits of of the mother liquor so the rate of evaporation crystal growth and the rapid evalution of vapour or rate of cooling or both must be progressively from the entering solution keeps the liquid in increased as the crystals grow in size.



Erg. 10.

the bottom of B sufficiently agitated that a certain amount of growth must occur before small crystals = wNd, *n. the crystals are capable of falling through the barometrie kep D and into the vessel E. All barometrie leg D and into the vessel E. the time that the crystals are descending the tube D, they remain in contact with the supersaturated solution. On reaching the supersaturated solution. On reaching the bottom of E the solution and crystals are trans ferred by the pump F to the settler G, and from thence to one or other of the centrifugal machines H for final separation of the crystals from the mother liquor. The overflow from the settler and the discharge from the centrifugals are returned to the vessel E, which is provided with an outlet K for mother liquor free from crystals.

OTREE METHODS.

Since crystal formation depends upon first, either the formation of nuclei or the addition of small seed crystals to the solution and, secondly, keeping the concentration of the solution at the working temperature in the metastable region if crystal size is the object or in the supersaturated region if fine crystals of varying sizes are desired, any method such as the adsorption of the solvent or the addition of some material to the mother liquor which diminishes the solubility of the substance, which it is desired to crystallise, in the solvent are means by which crystal formation can be effected. The addition of impurities to the mother liquor, however, even in small quantities affects materially both the rate of crystal formation and the rate of crystal growth.

As impurities inhibit the growth it is usual therefore in industrial practice to prefer one or other of the three methods outlined above as the principle upon which the crystalliser operates, rather than by introducing impurities to change the solubility of the dissolved material in the solvent,

CONDITIONS DUBING CRYSTAL GROWTH

If the rate at which the material is deposited on the faces of the crystal is to be kept constant, it is necessary to maintain the same concentration

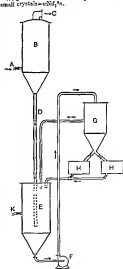
Let dimension of the amall crystals; d₂=mean dimension of the larger crystals grown from the smaller; n=number of faces on each crystal; N=number of crystals under consideration; S₁=total sur-fare of the small crystals; S₂=total surface of the large crystals; w=weight of material crystallised

per unit area of surface per unit time. With controlled crystallisation the crystals increase in size hut not in number, so that N is constant. The mean dimensions of the crystals are such that nd12 equals the surface of one of the small crystals and nd22 the ourface of the large crystal,

Total surface of small crystals $=S_1=Nd_1^2n$

Total surface of large crystals

 $=S_2=Nd_2^2n$. Weight of material removed per unit time by



Fra. 11.

large crystals= wNd_2^2n .

This shows that for a constant rate of deposition on the crystal surface much more material will be removed from the solution by large crystals than by small crystals in the same time: therefore the rate of cooling or the rate of evaporation or both must be progressively increased with the size of the crystal if the concentration of the solution is not to fall rapidly to the saturation line when further growth becomes impossible.

S. G. M. U. CRYSTOLON (CARBORUNDUM) v. ABRASIVES

CUBANITE. A copper ore with the composition CuFe₂S₃ (Cu 23·4%). It is usually massive with a brass-yellow or bronze-yellow colour and intimately intergrown with chalcopyritc, pyrrhotine, etc. Sp.gr. 4-1, hardness 3½. The mineral was first described by A. Breithaupt in 1843 from Barracanao in Cuba (hence the name), and soon afterwards recognised from Tunaberg in Sweden. It was long regarded as a rare mineral, supposed to be cubic in crystallisation, until polished sections of ores came to be examined by the metallographic method, when it was found to be of wide distribution. It was also found to be aniso-tropic, and therefore not cubic, and was identified with the small orthorhombic crystals of the same composition from the Morro Velho gold mine in Brazil, which E. Hussak in 1902 named chalmersite. Larger crystals have since been found at Sudbury in Canada (M. A. Peacock and G. M. Yatsevitch, Amer. Min. 1936, 21, 55).

L. J. S. CUBEBOL. The sesquiterpene alcohol, cubebol, C₁₅H₂₆O, m.p. 61-62°, occurs in oil of cubebs (Henderson and Robertson, J.C.S. 1930, 1908). It yields a phenylurethane, m.p. 186° and | percentage of dry matter):

Weight of material removed per unit time by an a-naphthylurethane, m.p. 197-198.5°. Tts structure has not been determined and it does not appear to be identical with cubeb-camphor examined by Blancbet and Sell, (Annalen, 1833, 6, 294).

J. L. S. CUBEBS, ESSENTIAL OIL OF. oil distilled from the dried, full-grown, unripe fruits of Piper cubeba Linn. (Fam. Piperaceæ), a tree indigenous to the Malay Archipelago, chiefly Java and Singapore; yield 10-15%. Other piperaceous fruits resembling cubebs are sometimes found in commerce.

Constituents.-Pinene, dipentene, and cadinene have been detected, and a sesquiterpene alcohol cubeb-camphor (m.p. 67°), which is sometimes separated from old oils. The blue colour is probably due to the presence of

azulene.

Characters.—A greenish or greenish-blue oil, sp.gr. 0.910 to 0.930, optical rotation -25° to -40° , ref. ind. 1.486-1.500. Soluble in 10 volumes of 90% alcohol.

C. T. B. Cucumis CUCUMBER. The fruit of Linn. The composition of cucumber varies particularly with size and with growth conditions, notably the level of supply of nitrogen to the roots (Dearborn, Cornell Agric. Exp. Sta. Mem. 1936, No. 192). Heinze (Z. Nahr.-Genussm. 1903, 6, 529, 577) records the following analyses:

Glu- Cane carbohy. Water, tein. Fat. cose, sugar, drates, Fibre, Ash. % % % % % % % % % % % Small 96·6 0·81 0·09 0·00 0·10 1·44 0·58 0·34 Large 95.8 0.67 0.09 0.66 0.09 1.60 1.65 0.42

Dearborn (l.c.) shows the composition of the whole plants on a somewhat different basis (as

	Stems.	Leaves.	Fruits.
	%	%	%
Soluble solids	23.6-26.7	15.2-22.8	58.9-58.3
Soluble N	0.212 - 0.562	0.151-0.365	0.304-0.710
Insoluble N	0.794-1.349	1.50-2.97	0.961-0.994
Amino-N	0.050-0.098	0.025-0.073	0.102 - 0.245
Amide-N	0.019 - 0.020	0.011-0.020	0.030-0.049
Nitrate-N	0.026 - 0.250	0.018-0.077	0.034-0.021
Lipin-N	0.003-0.000	0.027-0.064	0.004-0.041
Residual N	0.014 - 0.193	0.074-0.139	0.133 - 0.252
Reducing sugars	8.12-4.15	1.36-1.95	47.9-45.1
Non-reducing sugars	1.10-1.23	0.12-0.10	0.29-0.00
Starch	6.81-4.58	2.25-0.34	1.61 - 1.72
Acid-bydrolysable polysaccbarides .	2-99-2-89	12.01-4.81	2.79-2.65

Of the two figures given for each value the first refers to plants grown with a restricted, and the second to those grown with a generous, supply of nitrogen in the soil. According to M. Kitahara (J. Agric. Chem. Soc. Japan, 1936, 12, 595) the principal sugars are glucose and fructose. In hot water extracts of the fruit Yoshimura and Nishida (J. Chem. Soc. Japan, arginine.

Analysis of cucumber sceds is reported by Einhorn et al. (Maslob. Shir. Delo. 1929, 45, 44) as water 8.0, protein 29.7, fat 31.5, sugars, starch, etc. 1.9, pentosans 4.7, cellulose 13.9, pectins 0.6, phytin 1.1, lecithin 2.6, ash 3.9, P₂O₅ 2.25%. The oil contains fatty acids, comfructose. In hot water extracts of the fruit Prising stearic acid 3.72, palmitic acid 6.8, Yoshimura and Nishida (J. Chem. Soc. Japan, oleic acid 55.5, linolic acid 22.3%, and exbibits 1924, 45, 49) found adonine, trigonelline, and the following characteristics: d_{15} 0.9251, ref. ind. 1.4761 (25°), solidifying pt. -3.5°

456 CUCUMBER.

saponification value 191-1, iodine value 115 3, 0 87. acetyl value 16 6. acid value 3 5-10 7.

varies considerably with the manner and locality in which they are grown. The following data are representative values:

Small proportions of manganese, copper, and zine are also recorded by various analysts

CUDBEAR. A purplish red powder em ployed as a colouring agent, obtained by digesting Rocella lichen with ammonia

CULLEN EARTH or COLOGNE EARTH. A variety of brown coal or lignite,

in coal tar oil

Propylbenzanes — (1) Normal Propyl-BFIZENE (n cumone), C.H. CH, CH, CH, n Propylhenzene may be obtained by the general methods applicable to the formation of alkyl benzenes It is conveniently prepared (70-75%, yield) by the action of declays subjuste changed into the sopropyl, group owing to the on an othered solition of PRCH, Cl (* Organic Experiments. *New York, 195.*, tol. 4, p. 59.*).

Commerce has bp. 152 6-152 8/1700 mm, 4.*

Allyl cycloherane with platinised charcoal at 200°C. is converted to a mixture of propylcyclo hexane and propylbonzene (Levina and Tzurikov, J Gen. Chem. Russ 1934, 4, 1250). The equilibrium:

in the presence of a palladium catalyst has been investigated (ibid. 1933, 3, 718)

Propylbenzene (b p. 157°) has normal aromatic reactivity. Also it is interesting that it reacts with carbon tetrabromide giving C₆H₅·CHBr C₂H₆ (Hunter and Edgar, J Amer. Chem. Soc 1052, 54, 2025), and with methyl actone (von Miller and Robde, Ber. 1890. 23, 1070). Vigorous oxidation converts propyl the usual way (Gibson, J.CS. 1920, 117, 948, benzent to benzoie acidi. A number of pares Bert and Dorrer, Bell Soc.chim. 1927, 44, 1170) on the nitration of alkyl benzenes stodied from Chloranton of comeno jelds mono, di tribe point of view of modern electronic theories tetra, and penta chloro cumens and the have recently appeared in the Journal of corresponding chlorobenzenes. The isopropy the Chemical Society: p isopropyltolucne-Le group is thus easily displaced On nitration of Tevre, 1033, 980; p-etbyltolnene-Brady and Day, 1934, 114; n propylbenzene—Brady and Cunungham, 1934, 121; p tert butyltoluene—

Brady and Lahmi, 1934, 1934.

(2) 150 PROPYLBENZENE (CUMENE), - Forma. Reichert-Meissl value 0 5-1 05, Polenske value iton. Cumene may be synthesised by many general methods It may be prepared by the The composition of the ash of cucumbers action of stopropyl sulphate on phenyl magnessum bromide (Bert, Compt. rend. 1923, 176 840), by condensation of isopropyl alcohol and benzene in the presence of concentrated sulphung neid (Meyer, Monatsch. 1929, 53, 54, 721) and by condensation of stopropyl chloride and benzene in the presence of a little aluminium chloride (Radziewanowski, Ber., 1895, 28 1137). It is also readily obtained by boiling I part of p cymene, 10 parts of benzene, and 0 05 part of aluminium chloride for 10 hours, washing with water and fractionating (Bodtker and Halse, Bull. Soc. Chim. 1916, [iv], 19, 447):

$$C_6H_6M_6Pr^{\beta}+C_8H_6\rightarrow C_6H_5M_6+C_6H_5Pr^{\beta}$$

Cumens is formed with other propyl benzenes by the alkylation of benzene with propylene using various catalysts, e.g. H₂SO₄ or H₃PO₄ (USP. 2006541) or BF₂ (Nieuwland et al. J. Amer Chem Soc. 1935, 59, 1547) Cumene is EAHTH. A valuey of some comment of the produced when the vapours or certain response of limited use as a pagment. The name cument, are passed over heated metals [Ni, Cyl] (Sabatur organish) graven to septony) benzene to indicate et al. Compt rend 1919, 188, 871; 1919, 186 connection with cume read, from which it [190] (190), 2019, 201 frequently used as a genero term including (Mabery and Dunn, Amer Chem J. 1898, any of the isomero bearens hydrocarbons of the 18, 215). The preparation of cumen from formula C.H., Theory predicts the existence propyl bronde, bearens, and alumnum chor of eight of these, all of which have been found ide, and that of cymene from p bromotoluene, normal propyl bromide, and sodium (1:ttig, Schäfer, and König, Annalen, 1869, 149, 331) was for a long time the cause of great confusion in ascertaining the constitution of members of the cumene and cymene groups, as it was not realised that in each case the propyle group

0 862. With test butyl alcohol and concentrated sulphune acid cumrne yields ter butylcumens (H Barhier, Isième Congrès de Climne Industri-elle, Paris, Oct 1934). Vigorous oxidation of cumens yields bonzou acid. With chromic acid or permanganate in glacual sectic acid, phenyl dimethyl carbinol is produced (Brediter, Bull, See, Chim. 1991, [va], 25, 848). Cumena reacts with chromyl chloride in carbon dianiphide solution, giving an addition compound which on decomposition with water gives hydratropic aldehyde, C.H. CHMe CHO, and acetophenono (von Miller and Rohde, Ber. 1891, 24, 1357). Dilute nitric acid (d 1 075) in a sealed tube at 105°C. reacts with cument, giving βchromyl chloride, giving a double compound nitro \$\beta\$ phenylpropane (Nonowalou, Ber. 1894, that on decomposition with water yields benzyl 27, 468; 1895, 28, 1856).

Cumeno may be nutrated and sulphonated in tri, tetra, and penta chlorocumenes, the 150 propyl group is replaced by the nitro group (Qvist and Salo, Chem. Zentr. 1934, 2, 594). Propyl chloride, teri-butylbenzene, and p-di-teri-

butyl chloride and a very little aluminium chloride at room temperature (Bædtker, Bull. in sulphuric acid and crystallises out. On Soc. chim. 1906, [iii], 35, 834).

(3) Methylethylbenzenes.—The methylethylbenzenes, prepared by the action of sodium on a mixture of ethyl bromide or iodide with the corresponding bromo-toluenes, are liquids with boiling-points—o-, 158-159°, 158-159°, mp- 161-2°, and specific gravities of approximately 0.87 at 16°C.

Trimethylbenzenes.—(1) Consecutive trimethylbenzene (hemimellitene, hemimellitrine), $C_6H_3(CH_3)_3$ (1:2:3) is obtained by heating a-isoduric acid $C_6H_2(CH_3)_3CO_2H$ with lime. It has recently been isolated from an Oklahoma petrol (Beveridge and Sehicktanz, J. Res. Nat. Bur. Stand. 1933, 2, 665). The benzenoid hydrocarbons of the fraction, b.p. 118-132°/ 215 mm., were separated from other hydrocarbons by extraction with liquid sulphur dioxide and from one another by processes involving distillation, crystallisation, selective sulphonation, crystallisation of the sulphonic acids, and selective hydrolysis of the sulphonic acids. It was estimated that mesitylene, pseudo. cumene and hemimellitene are present in the crude petroleum used in the proportions 0.02, 0.2, and 0.06% respectively.

Hemimellitine has b.p. 176.1, sp.gr. 0.8951 $(20^{\circ}\text{C.}), n_{D}^{25} = 1.5116.$

(2) Unsymmetrical trimethylbenzene (ψ-cumene), C₆H₃(CH₃)₃ (1:2:4), occurs together with mesitylene in coal-tar (Beilstein and Kögler, Annalen, 1866, 137, 317), and is present in many petroleums (Poni, Chem. Zentr. 1906; 1, 459). It is prepared by the action of sodium on a mixture of either bromo-paraxylene or bromomelaxylene with methyl iodide. In order to metaylene with metayl rother. In other to isolate it from coal tar, Armstrong recommends (Chem. News, 1878, 38, 5; Bcr. 1878, 11, 1697) that the mixture of ψ -cumene and mesitylene (v. infra) boiling between 160° and 170°, obtained from coal tar by fractional distillation, should be converted into their sulphonic acids, and the mixture heated with concentrated hydrochloric acid in a sealed tube for 1 hour at 100°. In this way the mesitylene-sulphonic acid is decomposed, regenerating mesitylene, whilst the \(\psi\)-cumenesulphonic acid is not attacked. After separating the mesitylene the ψ -cumenesulphonic acid is precipitated from the hydrochloric acid solution by sulphuric acid, purified by recrystallisation from dilute sulphuric acid, and finally decomposed by heating with hydrochloric acid at $140-150^{\circ}$, when it breaks up into ψ -cumene and sulphuric acid. A slight modification of this method of scparation is described by Smith and Cass, J. Amer. Chem. Soc. 1932, 54, 1603). Jacobsen (Ber. 1876, 9, 256) separates the two isomeric sulphonamides by recrystallisation from alcohol, in which mesitylenesulphonamide is readily, ψ-cumencsulphonamide only sparingly, soluble. The sulphonamides are then hydrolysed by heating with concentrated hydrochloric acid at

Another method proposed by Jacobsen (Annalen, 1877, 184, 199) is to separate the two

butylbenzene are formed from cumene, iso-|sulphonic acids by fractional crystallisation; ψ -cumenesulphonic acid is soluble with difficulty heating to its melting point (110°) and passing in steam at a temperature of 250°, pure ψ-cumene is obtained (see also Schultz and Herzfeld, Ber. 1909, 42, 3602). ψ-Cumene boils at 168·2°, and has sp.gr. 0.877 at 18° (Schultz and Herzfeld, l.c.). On oxidation with nitric acid, it yields two isomeric acids of the formula C6H3(CH3)2CO2H-xylic acid and p-xylic acid-and a small quantity of the isomeric methylisophthalic and phthalic acids:

$C_6H_3(CH_3)(CO_2H)_2$

(Bentley and Perkin, J.C.S. 1897, 71, 165).

Oxidation of \(\psi\)-cumene with lead dioxide or manganese dioxide in the presence of sulphuric acid or electrolytically yields considerable quantities of 2:4-dimethylbenzaldehyde (Perkin and Law, J.C.S. 1907, 91, 263, 752). When boiled with aluminium chloride ψ -cumene is converted to a mixture of benzene, toluene, xylene, mesitylene, durene, etc. (Jacobsen, Ber. 1885, 18, 341; Anschütz, Annalen, 1886, 235, 186). This is a general type of reaction for polymethylbenzenes.

(3) Symmetrical Trimethylbenzene (mesitylene), C₆H₃(CH₃)₃ (1:3:5).—Mesitylene occurs in the coal tar fraction of b.p. 157-162° (Fittig and Wackenroder, Annalen, 1869, 151, 292; Schultz, Ber. 1909, 42, 3602), and in American petroleum. It is obtained on elimination of 3 molecules of water from 3 molecules of acetone. This elimination may be brought about by means of sulphuric acid (Organic Syntheses, 2,

Mesitylene is formed on dissolving allylene, CH3.C:CH in sulphuric acid and distilling with water (Fittig and Schrohe, Ber. 1875, 8, 17). Mesitylene has b.p. 164.5, sp.gr. 0.8768 4°/4°. It has the normal reactivity of a compound of its constitution.

CUMIDINES, C₉H₁₃N. The term "cumidine," which originally was proposed for the monobasic derivatives of cumene (isopropylbenzene) includes also by accepted usage numerous basic derivatives of pseudo- or (4)-cumene (1:2:4-trimethylbenzene, mesitylene (sym. trimethylbenzene), n-propylbenzene, and otber bases of the same empirical formula. Of these isomers only those of practical interest are treated below, the most important being sym- ψ cumidine and mesidine.

orthoCumidine, o-aminoisopropylbenzone, is formed by decarboxylation of aminocuminic acid with baryta (Cahours, Annalen, 1859,109, 19) and to a smaller extent by reduction of the crude nitration product of cumene. It yields normal salts and has b.p. 213.5-214.5/732 mm. Passage over hot lead peroxide yields indole. The acetyl derivative has m.p. 72°.

Cumidine, p-aminoisopropylbenzene, is made by nitration of cumene with fuming nitric acid and reduction of the nitrocompound with ammonium sulphide (Constam and Goldschmidt. Ber. 1888, 21, 1157) and also by direct condensation of aniline with isopropyl alcohol in presence of zinc chloride at 260° (Louis, Ber.

1883, 16, 111). Bp. 216-218°, acetyl deriva- fruit reveals four dorsal vittæ, or essential oil two mp. 102-102 5°. On oxidation with ducts, and an oily endosperm. The flavour and sodium chlorate and potassium ferrocyanide in hydrochlorie acid solution, cumiding yields an hit are more bitter and less agreeable. It is used "amilne-black" dyestuff (Kirpitschnikow, as a constituent of curry powder and as a Chem. Zentr. 1906, 1, 829) and it is also the carminative, the latter chiefly in retengary source of a dyestuff obtained by condensation with cyanogen chloride in pyridine solution (G.P. 155782).

baraAminopropyibenzene is formed by condensation of aniline with propyl alcohol (Willgerodt and Sckerl, Annalen, 1903, 327, 301) and has h p. 224-226°, acetyl derivative, m.p. 96° (for preparation, see G. Baddeley and J. Kenner, J.C S. 1935, 303).

Mesidine, obtained by reducing nitromesitylene with tin and acid (Fittig and Storer, Annalen, 1868, 147, 3) as well as hy numerous less usual methods of introducing an amino group, has hp. 229°, acetyl derivative, m.p. 216° Mesidino undergoes normal reactions and finds some use as a dye intermediate.

is obtained by reducing nitropseudocumene with tin and hydrochloric acid, but is more economically prepared by digesting technical xylidine with methyl alcohol at 300°, the basic mixture being separated by crystallisation of the nitrates (G.P. 22265). \$\psi\$ Cumidine has m.p. 63°, b p.234°, is mealuble in water and is characterised by a sparingly soluble nitrate and hy complexes with sym.truntrobenzene and sym truntro-tolusne (Noelting and Sommerhoff, Ber. 1906, 89, 78); acetyl derivative, m.p. 161°, ψ-cumidine while exhibiting many normal reactions undergoes some remarkable condensations and oxidations to substituted acridines (Senier and Compton, J.C.S. 1907, 94, 1934) and hydro quinolines (Jones and White, shid 1919, 97, 643) Reaction with a bromossovaleryl bromide yields a hypnotic Nalkyl derivative, but & cumiding is more important as the source of azo colours of the Ponceau senes (Schultz, Tab. No. 83).

Asym. / cumidine.

is prepared by reduction of the corresponding nitro-compound with iron and acetic acid (Edler, Ber. 1885, 18, 630), or with iron and aulphuric acid (Huender, Rec. trav. chim. 1915, 34, II), m.p. 36°, h.p. 233°, soluble in water.
CUMIN or CUMMIN. The dried fruit

(seed) of Cuminum Cyminum Linn, (Fam. Um bellifera), a small annual plant indigenous to Egypt, but cultivated in other parts of Northern Africa, Southern Europe and India. Tha fruits are about 4 to 6 mm. long, tapezing towards both base and apex, resembling caraways and, in fact, they have been called Roman caraways. The mericarps are less curved than those of caraway, and whilst they are usually separate, they may be attached to the pedicel. Each mericarp has five longitudinal ridges densely covered with numerous bairs, alternated with secondary ridges. The cross-section of the CUMMIN v. CUMIN.

odour are reminiscent of both caraway and anise. practice.

Mecroscopic Appearance.-The hairs are seen to be part of the pericarp, measuring up to 200n m length and from 25 to 40 u in breadth, and are composed of elongated cells. These are dis tinctive of the fruit. The vittee on the outer surfaces measure up to 200 µ in diameter, whilst those of the commissural side are even larger. Each of the primary ribs contains a bundle of fibres about 50μ in diameter. The endosperm contains fixed oil and alcurone grains which are 15µ or less in diameter and enclose rosettes of calcium oxalate.

Chemical Composition .- C. Arragon (Ann. Falsif. 1915, 8, 345) quotes the following composition of a sample of cumin fruit from Holland .

Water					10 5
Ash					7.27
Fat .					22 9
Esscutia	Ιo	ıl			23
Protein					226
Fibre					134

All figures, except water, are on the dry material

The principal constituent is the essential oil. and probably the most satisfactory method of determining this is by the process of Cocking and Middleton (Quart, Pharm. 1935, 8, 435). The powdered fruit is mixed with brine and distilled, the vapours being passed through the top of a condenser into a graduated tube, in which the oil is collected whilst the condensed water as returned to the distillation flock, An air inlet and outlet are provided by a side tube below the condenser, but above the graduated tube. Cumin contains from 25 to

4 0% of essential oil (q v.), Standards.—The British Pharmaceutical Codex " suggests that not more than 2% of foreign organic matter should be present.

CUMIN. ESSENTIAL OIL OF. oil distilled from the fruits of Cuminum Cyminum Linn. (Fam. Umbelliferæ), a native of Egypt and the Mediterranean, and cultivated in India and China. The chief supplies are from Morocco, Malta, and Sielly; yield 2 5 to 40%. Composition.—The chief constituent is cumic

T. McL.

or cuminic aldebyde, of which 30 to 40% is present. Pinene, cymene, dipentene, β phel-

landrene are present.

Characters—A colourless or pale yellow oil darkening on keeping. Sp gr. 0 900-0 935, opt. rotation +3° to +8°, ref. ind. 1 405-1 509. Solahlo in 10 volumes 80% alcohol. The cumic aldehyde is determined by the hydroxylamine process (t. ALDERYDES). C. T. B.

CUPFERRON. Ammonium-nitroso-Bphenylhydroxylamine. Used as a reagent for Cu, and also in the separation of Fe, Ti, and Zr v. Chemical Analysis, Vol. II, p. 583.

CUPRASE. Colloidal cupric hydroxide.

B-isoCUPREIDINE (isoaboQUINIDINE) v. CINCHONA ALKALOIDS (this vol., p. 170d).

CUPREINE v. CIPCHONA ALKALOIDS (this vol., p. 160d).

CUPRENE v. ACETYLENE.

CUPRETENINE v. CINCHONA ALKALOIDS (this vol., p. 178b).

CUPRI-ADEPTOL. Copper m-phenol sulphonate.

CUPRITE. Native cuprous oxide, Cu,O, crystallising in the cubic system. Brilliant, transparent crystals of a ruhy-red colour (hence the name ruby-copper-ore) are not uncommon; sp.gr. 6.0. In the variety called *chalcotrichite*, the crystals are capillary in form, being enormously elongated in the direction of one of the cuhic edges, and are loosely matted together in plush-like masses. Another variety, known as tile-ore (Ger. Ziegelerz), is earthy or compact and of a hrick-red or brownish colour; it consists of an intimate mixture of cuprite and limonite, and has resulted from the alteration of chalcopyrite. Cuprite is sometimes of importance as an ore in the upper oxidised portions of copper veins.

L. J. S.

CUPROL. Copper salt of nucleinic acid. External astringent dressing (Parke Davis, London). B.P.C. 1934.

CUPRON, a-benzoin oxime, used in the quantitative determination of copper and molybdenum v. Chemical Analysis, Vol. II, p. 605d.

CUPROSILICON. Cuprous silicide (v. COPPER).

CUPRUM LAZUREUM v. AZURITE.

CURACIT-SODA. A mixture of the sodium salts of glycocholic and other acids obtained from ox gall, used as a wetting-out

CURARE. Other native names, Curara, Urari, Woorari, Woorali, Woorara. Curare is the name given to the arrow poisons of S. American Indians who dwell in the region of the Orinoco and Amazon rivers. Knowledge of it was first hrought to Europe hy Sir Walter Raleigh in 1595, and from that time to the present its origin and preparation have heen discussed by explorers, hotanists and chemists. A summary of the earlier observations of La Condamine, Humboldt and Bonpland, Waterton, Schomhurgk, Castelnau, Thirion, Johert and Crévaux is given by Planchon (Pharm. J. decoction of several plants, one of which is usually a species of Strychnos, to which the toxic action of the preparation is due. The remainder contribute extractives which give the curare the used in the preparation of the product vary with the locality as the following shows:

Sources of Curare.—(1) Upper Amazon: Strychnos Castelnaeana, Baill., S. Yapurensis, Planch.

(2) Upper Orinoco: S. Gubleri, Planch.

(3) British Guiana: S. toxifera, Schomh.; S. Schomburgkiana, Klotzsch.; Benth.

(4) Upper French Guiana: S. Crevauxii, Planch.

(5) Esmeralda (Venezuela): S. guianensis, Aubl.

(6) Peruvian Amazon: Chondrodendron spp. (?).

The advent of firearms replaced arrows for hunting, and curare rarely appears nowadays on the European market. Such curare as is found pharmaceutical houses, and museums, physiological lahoratories consists of a hrown to hlack, often brittle, extract with a hitter taste. In medicine it has been used in the treatment of tetanus, hydrophohia, epilepsy, and in cases of chronic nervous disease with predominant muscular rigidity. For a summary, see West (Proc. Roy. Soc. Med. 1935, 28, 41). As native preparations vary considerably in composition the greatest care is necessary in the administration of the drug and its efficacy must first be tested on animals.

For the earlier chemical investigations on curare, see Flückiger (Arch. Pharm. 1890, 228, 78). The foundation of our chemical knowledge was laid by Boehm (Chem. Zentr. 1895, 2, 1084; 1897, 2, 1078; Arch. Pharm. 1897, 235, 660) who examined the three kinds of curarc, distinguished by their containers, which have at various times appeared in some quantity on the European market. Boehm showed that the highly active principle in each case was a different quaternary alkaloid. Tubocurare or para curare was imported in bamboo tubes and contained a crystalline inactive tertiary alkaloid curine which was shown later hy Spath, Leithe, and Ladeck (Ber. 1928, 61, [B], 1698) to he the lævo-enantiomorph of d-bebeerine, an alkaloid found in Radix Pareiræ bravæ (Chondrodendron tomentosum, R. and P.). Its formula is $C_{36}H_{38}O_6N_2$ (Späth and Kuffner, Ber. 1934, 67, [B], 55) and the structure of its di-O-methyl ether is now known in detail. It helongs to the group of hisbenzylisoquinoline alkaloids which are formed by fusion of two henzylisoquinoline structures through ether linkages (Faltis, Kadiera, and Dohlhammer, Ber. 1936, 69, [B], King, J.C.S. 1936, 1276). amorphous quaternary alkaloid called tubocurarine hy Boehm, was crystallised hy King (J.C.S. 1935, 1381) and found to produce complete "curare" paralysis of the frog on a dose of 0.5 mg. per kg. within 15 minutes. It was also 1880, [iii], 11, 469, 491, 529, 589, 693, 754). shown (King, l.c.) that d-tubocurarine chloride It is now known that curare is an aqueous had the properties of a phenolic betaine and on complete methylation of the phenolic groups gave d-O-methyltuhocurarine chloride which proved to he a diastereoisomeride of d-Omethylbebeerine methochloride (I). It is very consistency necessary for preservation of the probable that tubocurare is prepared from some poison and for tipping the arrows. The plants species of Chondrodendron hitherto not examined chemically and growing in the region of the Peruvian Amazon.

O-Methyltobocurarine chieride O-Methylbebeerine methochloride.

to contain an amorphous quaternary alkaloid, to as such as 3,000 tons in 1930 and about 2,000 C₁₁H₁₂ON₁Cl, called curarine, together with tons in succeeding years (cf. J. Cunha da taces of a non quaternary alkaloid. The same [Sulveira, Anasi do Instituto superior de agroamorphous quaternary alkaloid was isolated from Strychnos toxifera Schomb (King, Nature, 1935, 135, 469) thus confirming Robert Schom burgk'a discovery of S. toxifera as the main sources of the active ingredient of the curare of the Macus Indians from British Guiana It paralysed frogs on a dose of 0 125 mg per kg and showed colour reactions reminiscent of strychnine,

The third variety, known as pot curare, comes from Brazil, and appeared on the market in small jars usually of unglazed clay. Boehm found that this curare varied widely in activity but from ective material was able to isolate two relatively mactive non-quaternary alkaloids, protocurine, C20H22O2N, needles, mp. 306° (decomp), and protocuridine, prisms, mp. 274-276°, and an active amorphous quaternary base protocurarine of uncertain composition. The last-mentioned showed colour reactions similar to strychnine. The paralysing dose on frogs was 013 mg. per kg. (Jacabhazy, Areb. exp. Path. Pharm. 1899, 42, 10). It is probable that the active principle is contributed by Etrychnos Castelnacana Wedd, and not improbable that other species, such as Cocculus toxicoferus, Wedd, and Anomospermum grandsfolium Eichler supply some of the alkaloids present.

The predominant pharmacological action of principle in the three best known types of curare is a quaternary alkaloid is noteworthy since it is a pharmacological commonplace that nearly all ammonium bases paralyse motor nerve endings of voluntary muscle (Crum-Brown and Fraser, Proc. Roy. Soc. Edin. 1869, 6, 560).

The recent investigations of Freue in 8 America (Pharm. Ztg. 1933, 78, 852; 1938, 81, 818) show that a pot curare of the Chavante Indians of Southern Brazil is employed for killing animals to be used as food and is exported as an article of trade to the Argentine. It contains as its active principle a non quaternary alkaloid, macoubeine,

C,H,O,N,4H,O,

[a]D-555°, which is devoid of the typical "curare" action of the pharmacologist. For this reason it is better described as a "false curare." It is prepared from Macoubea Guyan. ensis Aubl. (Fam. Apocynaceæ).

H. K. CURARINE v. CUBARE.

CURCAS OIL (purging nut oil, physic nut oil, Oleum infernale; Fr. huile de pulghère,de purghère, - de gros pignon d'Inde), 13 obtained from the seeds of Jatropha cureas I., (Fam. Euphorhiaceæ), a shrub or small tree, prohably indigenous to Central America, but growing freely or under cultivation in many parts of the tropics, and especially in the Portuguese colonies. In the Cape Verde Islands, where the cultivation of cureas is second only to that of Calabash or gourd curare was found by Boehm | coffee, the export of seed to Lisbon smounted nomia (Lisbon), 1934, 6, fasc 1, 116); in Réunion the plant is grown as a support for the vanilla plant The seeds consist of from 32-40% of husk and 60-68% of kernel, containing from 46-58% of oil (i e about 30-40% on the whole seed) which is recovered commercially by hot expression The oil is pale in colour, becoming reddish vellow upon exposure to the air, and has a viscosity about equal to, or slightly greater than, that of olive oil: it is characterised by a disagreeable odour, and possesses emetic and atrongly purgative properties, which are much more pronounced than those of castor oil; it is distinguished from the latter by its lack of optical activity, low acetyl value and viscosity, and by its sparing solubility in alcohol and ready miscibility with light petroleum. It differs

or vesicant properties. The recorded analytical values for curess oil show considerable variations (probably depending partly upon the origin of the seed), but the following ranges are typical for genuine expressed oils: f. pt. c. -13°C., spgr. 15 0 918-0 921, acid value 07 to 92, saponification value 190-103, iodine value 93-100, acetyl value 5-10 (figures of 18 and 25 were recorded for two samples examined at the Imperial Institute (Bull. Imp. Inst. 1921, 19, 288). Samples of oil curare consists in a paralysis of the nerve endings (with 1mb), 1mb, 1921, 19, 285). Samples see of violuntary mucle, and for this reason it as exceeded perfection print from decordiscted much used in physiological research. The 1002, Françous and Droit, Bull. Soc. chim. 1933, theorem y. D. Bochm that the "curare" (w), 55, 782, 4 Afranta seed scritteded in the cold), (av), 53, 728; African seeds extracted in the cold), and by Adragens (IVe Congr. Intern. Tech. Chim.

from eroton oil (q v.) in possessing no rubefacient

1 A mixture of castor oil and cureas oil, sold under the name of "hagish castor oil" is stated to have been a popular purgative in America and the Antilles in the last century (see Diss, Ann. Inst. Col. Marsellies, 1896, 3, 26)

Ind. Agric., Bruxelles, 1935; Bull. Mat. Grasses, | 1936, 10, 813; seeds from Belgian Congo), had sp.gr. 15 0.9122-0.9219, ref. ind. 15 1.4694-1.4730, acid value 0.4—7.8, saponification value 176—180 (Droit), 193—196 (Adriaens); iodine value 97—98 (Droit), 96.6—106.9 (Adriaens); acetyl value 4.5—8.5, unsaponifiable matter 0.1—1.15%, including a phytosterol and a resinous material (Droit).

Whilst the commercial oils do not crystallise on exposure to low temperatures, the oils extracted in the laboratory gradually deposit crystals which appear to consist chiefly of oleodistearin. According to the detailed examination by Droit, the fatty acids of curcas oil consist solely of stearic, palmitic, myristic, oleic and linolic acids; no trace of ricinoleic or other hydroxyacid (such as the "curcinoleic acid," the presence of which was reported by Siegel and other early investigators) could be found (cf. Klein, Z. angew. Chem. 1898, 11, 1012). The "isocetic" acid isolated by Bouis (Compt. rend. 1854, 39, 923) appears to have been a mixture of palmitic and myristic acids. The medicinal use of the oil is limited by the presence in it of a toxic principle (a sterol-resin ester?) which is associated with the alcohol-soluble fraction of the oil (cf. Cadet de Gassicourt, J. de Pharm. (Paris), 1824, 10, 170; Soubeiran, ibid. 1829, 15, 500; Felke, l.c., infra; Droit, l.c., infra), and is not destroyed by heating for 1½ hours at 100°C.; upon saponification, this fraction yields fatty acids, a phytosterol, and a resin, none of which appears to have any physiological action when examined separately (Droit, Bull. Mat. Grasses, 1932, 16, 270).

A second poisonous substance, curcine (a toxalbumin) (cf. Stillmark, Arb. Pharm. Inst. Dorpat, 1889, 3, 149; Siegel, Thesis, Dorpat, 1893; Felke, Landw. Versuchs-Stat. 1913, 82,427) is found in the decorticated seeds after removal of the oil, and, consequently, the by-product seed-cake can only be utilised as a fertiliser. Curcas seeds further contain a fairly active lipase (Grimme, Z. deut. Öl-Fett-Ind., 1921, 41, 51).

In addition to its use in medicine, curcas oil is used in the manufacture of soap (both in the tropics and in Europe), and to a limited extent as a burning oil; its use as a substitute for olive oil in compound lubricating oils has been suggested. The oil thickens somewhat on heating, and is frequently reported to possess weak drying properties (cf. Bull. Imp. Inst. 1921, 19, 288); this propensity appears to be less well marked, however, in the oils from the Congo (Adriaens, l.c.).

(For a detailed study of the purging nut, its oil and toxicology, and bibliography, see Droit, l.c.; also Anon., Gov. Lab. Bangkok, 1929, No. 4, 8).

E.L.

CURCINE v. CURCAS OIL.

CURCUMENES, THE. The essential oil from the rhizomes of Curcuma aromatica contains

monohydrochloride, b.p. $150-155^{\circ}/8$ mm., has b.p. $128-130^{\circ}/7$ mm., d_{30}^{30} 0.8633, $n_{\rm D}^{30}$ 1.4944, $[a]_{D}$ -22.9°. It contains three ethylenic linkages and is best characterised by the preparation of its nitrosate, m.p. 101°. l-β-Curcumene, which can be purified through its trihydrochloride, m.p. 84-85°, has b.p. 128-130°/6 mm., d_0^{30} 0.8810, $n_{\rm D}^{30}$ 1.4949, [a]_D -27.9°. It has three ethylenic linkages. The constitutions of these hydrocarbons have not been determined.

J. L. S. CURCUMIN (Turmeric yellow), the yellow substantive dye and indicator from the roots of Curcuma species Linn.

CURINE v. CURARE.

CURITE, 2PbO·5UO₃·4H₂O, a uranium mincral.

CURRANT. The fruit of bushy species of Ribes. Two varieties are commonly met, the red currant, R. rubrum (or R. vulgare Lam.), and the black currant, R. nigrum, L. Variants of R. rubrum in which the fruit is white are specially favoured for dessert.

Analyses of the red currants from American

and European sources respectively are:

Acids Total Pro-Invert Sucsolids. tein. citric). sugar. rose. Pectin. Ash. 2·21 (1) 12.97 3·44 1.37 0.0 0.6 (2) 15.291.90 5.740.460.57

Munson et al., U.S. Dept. Agric. Bur. Chem. Bull. 1905, No. 66.
 Olig, Z. Unters. Nahr. Genussm. 1910, 19, 558.

According to Hotter (Z. landw. Versuchsw. 1906, 9, 747) the proportion of fructose is slightly higher than that of glucose. The actual amounts of sucrose present in the fruit and the proportion of reducing to non-reducing sugars varies considerably with the stage of ripeness of the currant. Sobolevskaja and Turetzkaja (Bull. Acad. Sci. U.R.S.S. [Classe sci. math.], 1934, 9, 1341), have observed the steady increase in total sugars and in the ratio of invert sugar to sucrose as the fruit develops. The absolute amount of sucrose, however, increases markedly during the final stages of ripening. These authors explain the general course of sugar changes in the ripening fruit

 $sucrose \rightarrow glucose + fructose \rightarrow polysaccharide \rightarrow$ glucose+fructose.

The accumulation of sucrose in ripened fruit is ascribed to the cessation of its decomposition rather than to its re synthesis from invert sugar; the simultaneous increase in invert sugar is attributed to the hydrolysis of polysaccharides. The acidity of the fruit is due almost entirely to citric acid, although small amounts of malic acid are occasionally reported.

The seeds of red currants contain water 11.42, protein 14.92, fat 23.6, N-free extract 24.4. a mixture of two monocyclic sesquiterpene hydrocarbons C₁₅H₂₄, a- and β-curcumenes (Rao, Shintre and Simonsen, J. Indian Inst. Sci. 1926, 9, A, 140; Rao and Simonsen, J.C.S. 1928, 1926, 9, A, 140; Rao and Simonsen, J.C.S. 1928, ref. ind. 1·4772 (25°), saponification value 194·5, 2496). *l*-a-Curcumenc, purified through its iodine value 159·8, Reichert-Meissl value 0·55, value 12 9, unsaponifiable matter 0 64%.

The mineral constituents of red currents (in terms of fresh fruit) include: total ash 041.

 K.O. 0 19, Na.O 0 02, CaO 0 08, MgO 0 03,
 P.O. 0 09, Fe 0 0007, Al 0 0015, Zn 0 0002%.
 Windisch and Schmidt (Z. Unters. Nahr.-Genussm. 1909, 17, 584) record the following as average analyses of the juice of red currents: d₁₅ 1 045, solids 11 65, protein 0 34, acids (as citric) 2 11, invert sugar 6 90, total ash 0 48 g.

per 100 c c.

The black current resembles the red species in composition in many respects. Detailed analyses are, however, less common. Hotter (Le) records the following analysis for black currant pulp: solids 20 7-20 9, acids (as male) 23-3-4, total sngars 7-3-7 9, glucose 3 3-3 5, fractose 40-44, sucrose 02-04, tannin 033-041, ash 0 63-0 87.

The acids of the fruit comprise chiefly citine and tartane; the proportions vary with stage of ripening, that of citric acid reaching maximum in the half ripe stage.

As in the case of red currents the black variety undergo a very rapid increase in augar content in the last stages of ripening (or over ripening).

A. G. Po DEGRADATION OF CURTIUS ACID AZIDES v. AMIYES.

CUSCAMIDINE v. CINCHONA ALEALOIDS

(this vol., p. 161c). CUSCAMINE v. CINCHONA ALKALOIDS (this

vol , p. 161c). CUSCOHYGRINE v Cocaine and other

alkaloids of the Erythroxylon species. CUSCONIDINE v. CINCHONA ALEALOIDS (this vol , p. 161d).

CUSCONINE c. CINCHONA ALKALOIDS (this

vol., p. 161d).

CUSPARIA BARK. CUSPARINE, GALIPINE AND MINOR ALKALOIDS. Cusparia bark, Angostura bark, Cortex Cusparia: (Fr. Ecorce d'Angosture ; Ger Angosturarinde). It is denved from Galipea officinalis Hancock, ayn. Cusporta trifoliata Engler (Fam Rutacem), a small tree of Venezuela, and was originally shipped from Angostura on the Orinoco to Trinidad. It was long considered to be derived from the closely related Cusparia febrifuga D C. Brazilian (false) "Angostura" bark is derived from Esenbeckia febrifuga A. Juss. (Fam. Rutacex). For history and botanical investigation of different banks v. Hartwich and Gamper (Arch. Pharm. 1900, 238, 568)

Angostura bark contains about ten alkaloids, some of which are very simple basia compounds; it also contains a little essential oil,

a bitter principle and a gluroside.

The cusparia alkaloids were investigated by Tröger and Beckurts; for references concerning earlier investigations, v. Spath and Brunner

(Ber. 1924, 57, [B], 1243).

The best description for isolation and identifieation of cuspana alkaloids is given by Spath and Eberstaller (Ber. 1924, 57, [B], 1687), Spath and Papaioanou (Monatah, 1929, 52, 129), and and Papaioanou (Monatsh. 1029, 52, 129), and colourless oil with very faint odour, easily Spath and Phl (Ber. 1929, 62, IB), 2244, Monatsh, soluble in dilute HCI, B picrate, needles from 1930, 55, 3329, 16 kg, of aggostura bark MeOH, mp, 132°, B, phalinichloride forms

Polenske value 0-5, Hehner number 95 6, acid are percolated with EtOH for 8 days. EtOH is evaporated in vacuo and the only residue still containing appreciable amounts of EtOH (this is essential) diluted with 2 litres of 20% NaOH and extracted with 4 litres of Et.O. Phenolic and non-phenolic bases are thus scparated.

Non phenolic Bases .- By extracting above ethereal solution with 1% HCl the sparingly soluble bydrochlorides of cusparine (1) and galspine (11) are obtained. The two alkaloids are separated through the oxalates, a method developed by Tröger and Kroseberg (Arch. Pharm. 1912, 250, 503; (1) oxalate spaningly, (II)-oxalate easily, soluble in H₂O). Yields: (I)=1 06%, (II)=0 35%. 200 g of the oily mixture left after separation

of (I) and (II) are treated with light petroleum, which dissolves shout 30 g. The oily readue obtained by evaporation of the petroleum is steam distilled and the distillate re-distilled in vacuo, after a small first fraction A, 87 g. of

almost pure 2 n amai ares reaction A, 5 , 8, or almost pure 2 n amy 4 methoxy quinoline (III) distila at 190-200°/14 mm.; yield 0 05%.

The first fraction A is distilled again, when two separate fractions are obtained; bp. 100-145°/14 mm.=B (0.5 g) and bp. 145-190′/14 mm.=C (5.2 g.). Fraction B finally yields quinoline (1V) and 2 methylgainoline (quinaldine) (V). Yield of the two bases about 0 003%. Fraction C gives 1 methyl-2-keto 12-dhydrogumoine (VI) (0 01%) and 2 n amyl-qumoine (VII) (0 003%). Phenoic Bases.—Tho alkaline solution left

after asparation of the non-phenolic bases is acrdified with 5% HCI, separated from tar, basified with sods, and extracted with CHCI After evaporation the residue is extracted with ether and finally pure galipolina (VIII) is

obtained.

Cuspareine (Chem. Zentr. 1909, II, 1570; 1911, I, 163), galipoidine (Arch. Pharm. 1913, 251, 252), and angosturine (Chem. Zentr. 1911, 1, 163) were isolated by Troger and co workers, but their existence has not yet been confirmed. All cusparia alkaloids are optically mactive.

Cusparine (1), C1,H1,O1N, crystallises in needles from light petroleum, m.p. 92°, it is apparently trimorphous (colourless needles, yellow needles, and amber tinted crystals) (I) is easily acluble in organic solvents. The salts are sparingly soluble and may be readily separated from the salts of associated alkaloids B HCI+3H,O, white crystals; Boxalate is characteristic (v. supra), mp. 152-156, aulphur, yellow needles. Also platinichloride and aurichloride are crystelline. B methodide, m p. 176° (yellow prisms). (1) is the easiest of the angostura alkaloids to separate.

Galepine (II), C₂₀H₂₁O₃N, crystallises from EtOH, Et₂O or light petroleum in colourless prisms, m p. 1155°; it yields crystalline salts, which are more soluble than those of (I). B HBr, m p. 169° (yellow); B HI, m p. 178° (yellow); B methiodide, mp. 145° (yellow

needles).

2 n-Amyl 4 methoxy quinoline(III), C15 H15ON,

characteristic, vellow crystals, 220° m.p. (decomp.).

Quinoline (IV) and 2-methylquinoline (V) (quinaldine) were identified as picrates and 2:4:6-trinitro-1:3-cresolates.

1-Methyl-2-keto-1:2-dihydroquinoline (VI), C₁₀H₉ON, m.p. 74°; B-picrate from MeOH, m.p. 129-130°. 2-n-Amylquinoline (VII), h.p. 130-145°

10 mm., identified as picrate, m.p. 125-126°

(from MeOH).

Galipoline (VIII), C₁₉H₁₉O₃N, colourless crystals, m.p. 193° (from H₂O), soluble in caustic soda; by methylation of (VIII) with diazomethane (II) is obtained.

Cuspareine (cf. Tröger and Runne, Arch. Pharm. 1911, 249, 176), C₁₈H₁₉O₂N, m.p. 56°, is a very weak base and distils without decomposition. No salts could be obtained, it contains two MeO-groups. B·methiodide, m.p. 156°

(leaflets from H₂O).

(Galipoidine (cf. Tröger and Runne, l.c., 183),

C₁₉H₁₅O₄N, m.p. 233°, needles from EtOH,
insoluble in ligroin, light petroleum and benzene; sparingly soluble in hot EtOH. It
yields a crystalline platinichloride and aur chloride.

Angosturine is apparently identical with galipoidine (cf. Chem. Zentr. 1911, I, 164).

CONSTITUTION OF ANGOSTURA (CUSPARIA) ALKALOIDS.—When (I)-methiodide is treated with silver oxide or caustic soda, a new hase, isocusparine (m.p. 194°), is formed. (II)methiodide undergoes the same rearrangement. Apparenty the Me of the MeO-group migrates to the N-atom (cf. Tröger and Müller, Arch. Pharm. 1914, 252, 459), a rearrangement which is typical for a and γ-methoxy-quinolines (cf. dictamnine and skimmianine). By zinc dust distillation of cuspareine (Tröger and Runne, l.c., 182) and (II) (Tröger and Kroseherg, l.c., 525) quinoline was obtained. By potash-fusion hoth (1) and (II) yielded protocatechnic acid, on oxidation with permanganate veratric and a methoxyquinoline carhoxylic acid were obtained from (II). Tröger and Bönicke (Arch. Pharm. 1920, 258, 250) suggested a formula for (II) which was slightly modified by Späth and Brunner (l.c.); (I), (II), and (VIII) have the following formulæ, as proved by synthesis:

$$\begin{array}{c|c} R_{3} & & & R_{2} \\ \hline C > CH & & R_{2} \\ \hline C - CH_{2} - CH_{2} - CH_{2} \\ \hline (I): R_{1} + R_{2} = CH_{2} & G; \ R_{3} = OMe. \end{array}$$

(II): $R_1 = R_2 = R_3 = OMe$. (VIII): $R_1 = R_2 = OMe$; $R_3 = OH$.

Synthesis of (I): Spath and Brunner (l.c.). Synthesis of (II): Spath and Eberstaller (l.c.). Synthesis of (III): Spath and Pikl (l.c.).

Synthesis of (VIII): Spath and Papaioanou

Nothing is known about the constitution of cuspareine and galipoidine.

The essential oil (from Cusparia trifoliata) is described in Schimmel and Co.'s Semi-Annual Report, April, 1913.

Angosturin, C₀H₁₂O₅, m.p. 58°, is a crystalline bitter principle, readily soluble in H₂O and EtOH, but not in (Et)₂O. Angostura bark is a constituent of "Angostura bitters," and is used in medicine as a tonic in derangements of the alimentary canal, but it is not included in "British Pharmacopæia" nor in the "United States Pharmacopæia."

Schl.

CUSPARINE v. Cusparia Bark.

CUSSO, KOUSSO. The dried panicles of pistillate flowers of Brayera anthelmintica Kunth.

CUTCH v. CATECHU.

CYAMELIDE CYANIDES (this vol., p. 505).

CYANAMIDE v. CYANIDES (this vol., p. 505).

CYANANTHROL B (SOLWAY BLUE R) v. Anthraquinone Dyestuffs.

CYANIC ACID v. CYANIDES (this vol., p. 506d).

CYANIDATION **PROCESS** GOLD AND SILVER ORES v. CYANIDES. CYANIDES.

HISTORICAL.—The earliest known cyanide compound was Prussian blue, which was discovered accidentally in 1704 hy Dieshach and Dippel in an attempted preparation of Florentine lake. The new hlue very quickly displaced ultramarine, hut its method of manufacture remained a secret until 1724, when Woodward described the calcination of blood with potassium carhonate, and the treatment of the aqueous extract with ferrous sulphate and alum to give a greenish precipitate, which yielded Prussian blue when treated with hydrochloric acid. Macquer showed that horn, leather and other animal matter could he used instead of blood, and in 1752 found that Prussian blue was decomposed hy hoiling with alkali into ferric hydroxide and a soluble salt from which the blue could be regenerated. Potassium ferrocyanide (yellow prussiate) was first obtained in the crystalline form by Sage in 1772.

The relationships of the ferrocyanides, hydrocyanic acid, cyanogen and many of their derivatives were elucidated by the brilliant researches of Scheele (1782-3), Berthollet (1790), and Gay Lussac (1815). Scheele was the first to prepare an aqueous solution of hydrocyanic acid and Gay Lussac first isolated the anhydrous liquid.

Animal matter treated by the described ahove was the sole source of cyanides for about 150 years, until a new one was found in the cyanogen content of coal gas. The existence of cyanogen compounds in coal gas was mentioned by Jacquemyns in 1843, and in 1860 prussiate was manufactured from spent oxide for the first time at the Paris gas works. By 1884 gas works ferrocyanide had almost completely replaced the older product in the European industry, partly because of the

increased use of nitrogenous organic matter as a | was introduced in 1903, first in the citrus groves fertiliser, and partly because of the reduced of California to control a parasitic scale insect. consumption of Prussian blue due to the com-petition of coal tar dyes. In the United States and warehouses and other buildings against the old process held its own for nearly twenty yeard longer. Between 1890 and 1990, greater was formerly generated in a rist from cyanide and efficiency in the recovery of cyanides from coal acid, but since 1917 haud by drocyane acid has distillation products was obtained by the been marketed for direct application. introduction at gas works of special cyanide washers, the cyanide being almost quantitatively fixed as ferrocyanide which is relatively easily recovered from the scrubbing hourd.

The discovery of the electroplating process for gold and silver by J. R. and H. Elkington in solution of iron carbide and iron nitride in 1840 led to a demand for potassium cyamide, metallic iron. which was met by the process of fusing ferrocyanide with potassium carbonate, a reaction successful cyanide processes is ammonia or discovered in 1834 by F. and E. Rodgers methylamines (Schlempe process), but the Wis In 1887 MacAthur and the Forrests patential (1914-18) stimulated the search for methods of the cyanide process for the extraction of gold dixing atmospheric nitrogen as cyanide. Possor and silver from their ores, and it was first operated at Karangahake, New Zealand, in 1889, potassium cyanide from potassium carbonate, and near Johannesburg, Transvasl, in 1890 The cyanide process was rapidly adopted in all greatest advance towards the technical realisaand cyanue process was rapuny acopiect in an greace stavanice towards the technical reshina-tion of this reaction was made by Bucher and transformed the cyanule industry from a treatively large rapithetic nutienty. While the production, but the United States between 1912 and 1918, with the stimulus of shormal war conditions expithetic nutienty. While the production of potassium cyanules in 1889 (by the Redgers processed in the contract of the contract about the state of the contract about th process) was 50-70 tons per sinum, and of the same time Caro and Frank in Germany wers yellow prussate not more than 5,000 tons, it is endeavouring to produce cyanide by the estimated that the world production of potassium introgenation of calcium carbide, and the eyanide had reached 6,500 tons per annum hy 1899.

At first the increased quantities of eyanide were made from ferrocyanide by the Rodgers process, and then by the Edramoyer process now produced by this reaction. That no greater (discovered in 1870) using metalla sodium. The development of the fixation of introgen as rapidly increasing demand, however, stim-lated cyantle has occurred may be augusted to the search for direct synthetic processes, of the great progress which was made, both which a great number were invented and the during and immediately after the War, in those manifecture of synthetic ammounts, and the process of the auccess:

| Date of

Process	Location of plant.	first pro- duction
Siepermann	Stassfurt, Germany	1892
Beilby	Glasgow, Scotland	1892
British Cyan- ides Co.	Oldhury, Eogland	1895
Raschen	Runcorn, Eogland	1898
Readman	Leven, Scotland	1899
Castner	Germany, Scotland,	1899
	US.A.	1900
Bueb (Schlempe)	Dessau, Germany	1902
(Schlempc)	<u> </u>	<u> </u>

Of these the last two are still in operation and provide the greater part of the world's cyanide requirements. At the present time there is a tendency in the United States for a reversal many countries since 1931 has caused an of the old economic relationship, so that increased consumption by stimulating the goldaynthetic cyanide is the raw material for ferrocyanide.

The use of cyanide as a fumigant in peat control for case hardening and fumigation.

Another use of cyanide which has been developed since 1918 is in the case hardening of iron and steel, which now consumes increasing quantities of sodium cyanide. This use depends on the formation of a surface layer of a solid

The source of the nitrogen for the commercially and Boissière first attempted the manufacture of charcoal, and mirogen as early as 1843, but the process was brought to commercial success by Landis in America in 1918. Relatively large quantities of impure calcium cyanide, used in the gold mining industry and in fumigation, are which has now largely replaced by-product ammonia as the raw material for cyanide manu-

facture. The most recent research on eyanide synthesis has been in the realm of high temperature gas reactions for the production of hydrocyanic acid from ammonia (or nitrogen) and hydrocarbons, end on methods of converting the hydrocyanic acid so obtained into sodium cyanide of similar purity to the bigh-grade product of the Castner process. Of the many such processes described, only one, involving the passage of a gaseous mixture of nitrogen and hydrocarbon gas or vapour through an are, is known with certainty to have been operated on a commercial scale. It may well be, however, that the future development of the cyanide industry will be along these lines.

The world consumption of cyanide at the present time is probably more than 30,000 tons per year of sodium cyanide or its equivalent. The abandonment of the Gold Standard in mining end base metal industries, and steady development has occurred in the use of cyanide

under the following headings:

I. Complex iron cyanides (ferrocyanides, ferricyanides, pentacyanides).

Alkali metal cyanides.

III. Heavy metal cyanides (of copper, gold, mercury, silver, zinc).

IV. Alkaline earth cyanides. V. Hydrocyanic acid.

VI. Cyanogen and its derivatives.

VII. Cyanates.

VIII. Thiocyanates.

I. COMPLEX IRON CYANIDES.

GENERAL CHEMISTRY.-In this section a brief account is given of the chemical characteristics and relationships of the ferrocyanides

and ferricyanides.

In all probability the simple parent substances, ferrous cyanide, Fe(CN)2, and ferric cyanide, Fe(CN)₃, do not exist. Substances having the empirical composition of the expected ferrous cyanide are indeed produced in the dry way by heating ammonium ferrocyanide or hydroferrocyanic acid, and in the wet way by precipitating ferrous solutions with cyanide; but these are rather to be regarded as ferrous ferrocyanide, Fe₂[Fe(CN)₆], and similar complex derivatives of hydroferrocyanic acid, which are "polymers" of Fe(CN)₂ (see Wyrouboff, Ann. Chim. Phys. 1869, [iv], 16, 280; Hofmann, Arnoldi and Hiendlmaier, Annalen, 1907, 352, 54). Analogous preparations of Fe(CN)3 are lacking, for the thermal decomposition of ammonium ferricyanide and hydroferricyanic acid has not been studied, and only ferric hydroxido is produced by the interaction of cyanide with ferric salt solutions. A crystalline solid of the composition Fe₂(CN)₆,2H₂O has been prepared by Reihlen and von Kummer (Annalcn, 1929, 469, 30) from solutions of potassium ferricyanide and ferric nitrate, and this is most probably ferric ferricyanide,

Fe[Fe(CN)₆].

The iron hexacyanido ions are among the most stable complexes encountered in inorganic chemistry, the degree of dissociation into simpler ions being too small to be measured. It appears unlikely that a dissociation yielding the simple ferrous or ferric ions occurs at all, and when Fe++ or Fe+++ and CN' are brought together, formation of complex iron cyanide ions proceeds until one or other of the constituent ions has disappeared. This explains the failure of attempts to prepare the simple ferrous and ferric cyanides by wet reactions. The formation of ferrocyanides from cyanides occurs with insoluble as well as soluble ferrous compounds, and even with metallic iron itself. The

ionic equations

(a) $Fe^{+} + 6CN' = Fe(CN)_{6}''''$ (b) $FeS+6CN'=Fe(CN)_6''''+S''$

(c) $Fe(OH)_2 + 6CN' = Fe(CN)_6'''' + 2OH'$

(d) $Fe+2H_2O+6CN'$

 $= Fe(CN)_6'''' + 2OH' + H_2$

testify further to the extreme stability of the ferrocyanide ion, for if any appreciable dissocia-

The cyanide compounds are considered below | product of ferrous sulphide (b) or ferrous hydroxide (c) and (d) would be exceeded and the reaction reversed.

Ferricyanides cannot be synthesised from cyanides and ferric compounds by reactions analogous to those mentioned above, for cyanides can reduce ferric salts and ferric hydroxide, so that ferrocyanides result. According to Rupp (Z. anal. Chem. 1931, 86, 217), however, if the ferrie salt is first protected by conversion with ammonium phosphate into soluble complex ferric phosphate, quantitative formation of ferricyanide occurs on boiling with potassium cyanide. Ferricyanides are generally produced by oxidising ferrocyanides, either electrolytically or with chlorine or bromine. The transition from Fe(CN)₆"" to Fe(CN)₆" can be reversed under suitable conditions of experiment. For instance (X=Cl or Br)

 $2Fe(CN)_6''''+X_2 \rightleftharpoons 2Fe(CN)_6'''+2X'$

(Gillet, Bull. Soc. chim. Belg. 1912, 26, 236). In strongly acid solution under suitable conditions, potassium iodide is oxidised to iodine by ferricyanide, whilst in neutral solution ferroeyanide reduces iodine. Oxygen itself will oxidise ferrocyanide to ferricyanide in acid solution, but not in alkaline.

The commonest technical forms of ferrocyanide are "yellow prussiate of potasb," K4Fe(CN)6, and Prussian blue (ferric ferrocyanide, etc.). Other ferrocyanides can be obtained from the potassium salt by double decomposition with metallic salts, and from Prussian blue by boiling with the hydroxides or carbonates of tho alkali and alkaline carth metals, when solutions of the corresponding ferrocyanides are formed. Neither of these methods is entirely satisfactory for the production of pure compounds, for the products are usually contaminated by double salts which are formed with great case and are not easily removed. If pure products are required, it is necessary to neutralise hydroferrocyanic acid with the appropriate base, or to treat barium ferrocyanide with the appropriate sulphate.

The ferrocyanides and ferricyanides of the alkali and alkaline earth metals are soluble in water, generally crystallising with water of crystallisation. The ferrocyanides are yellowish in the hydrated form, and almost white when anhydrous. The ferricyanides have a characteristic ruby-red colour. The heavy metal salts of both anions are generally insoluble amorphous precipitates, the ferrocyanides having characteristic colours which are not related to those of the component ions (e.g. cupric ferrocyanide is brownish-red, ferric ferrocyanide is deep blue), and this is held to indicate that a constitutional change occurs on their formation.

When heated above 400°C, the solid ferrocyanides are decomposed into the constituent cyanides, e.g. KCN and "Fe(CN)₂," the latter decomposing further into iron, carbon, iron carbide and nitrogen. Where one of the cyanides is volatile (e.g. NH4CN or HCN), the primary decomposition, helped by the escape of the volatile product, occurs at a lower temperature. The solid ferricyanides undergo tion into ferrous ions occurred, the solubility similar disruption on heating, and are less

CYANIDES.

disruption of the iron cyanide complex also escapsog vapours as hydrocyanic acid, ammoni, occurs when the solid salts react with aulphurie and organic bases, and up to 57% of the nitrogen occurs with the such HCN is produced, and remaining in the obstred mass as converted with concentrated acid the product is carbon into ferrorganide, corresponding to an overall monoscide, derived from HCN by its hydrolysis to ammonia and formic acid, the latter being Although the reactions proceed equally well subsequently debydrated. In the case of ferricyanide, carbon dioxide also appears in the gaseous product, for the ferme salt arising from the initial disruption of the ferricyanide complex | Na Fe(CN), aq make the isolation of the solid can exidise some of the formic acid to carbon salt more difficult. dioxide and water.

Perrocyanide and ferricyanide solutions are electrolytically dissociated into metallic kations, that potassium cyanide is formed during the and iron cyanide complex amons. As already stated, the complex amons are very stable and give rise to no iron kations in solution Under certain conditions, bowever, they are subject | to a fission of one of the six cyanide groups, which is replaced by another group to form the complex pentacyanides. For instance, ferro evanide solutions, on heating alone, on treating with acids, or by the action of light in the absence of acids, become alkalino and evolve HCN, forming aquopentacyanides in solution. The reaction, which is characterised by a yellowing of the ferrocyanide solution, is ionically represented as follows:

Fe(CN),""+2H2O =[Fe(CN), H,O]"+OH+HCN

A similar reaction occurs, but more readily, with ferricyanide solutions; in the presence of nitro acid the seceding cyanide group may be replaced by NO to give nitrosopentacyanide ("nitroprusside"), and with carbon monoxide carbonylpentacyanide is produced.

Ferrocvanides.

TECHNICAL METHODS OF PREPARATION. From Nitrogenous Organic Matter .- The manufacture of ferrocyanides from nitrogenous organic matter by the process which led to the descovery of Prussian blue, and which was almost the sole source of ferrocyanide and cyanide until 1860, bas now been entirely superseded and is only of historical interest. The forms of nitrogenous matter used were dried blood, hore, hair, waste wool and feathers, or the animal charcoal obtained by their destructive distillation. The material was fused with potassium carbonate and iron turnings in east iron muliles or in rever beratory furnaces, and the melt was leached with bot water to obtain an aqueous solution from which potassium ferrocyanide erystallised out The black insoluble residue from the leaching contained double sulphides of potassium and ferrous and ferric iron, and various double silicates of potassium, calcium and aluminium The loss of potash in this residue appears to have been 45-90% of that usefully consumed,

The efficiency of ferrocyanide recovery in the process, based on the nitrogen content of the organio of the organic matter with potesh, not fusion of the organic matter with potesh, not more than 20%, of the ferroyanide theoretically Report, 1906, 68.

3. Chief Inspector of Alkah worss, more than 20%, of the ferroyanide theoretically Report, 1906, 68.

4. J. McLeod, J.S. C.I. 1907, 28, 137; Alkah 1007, 915. obtainable was actually recovered (Karmrodt, 4. J. McLood, J.S.C.I. 1907, Wagners Jahresber. 1857, 3, 139). When the Works, 44th Report, 1907, 215.

stable than the ferrocyanides, a loss of eyanogen introgenous matter was first carbonised alone, being detectable even at 180°C. Complete about 80% of the nitrogen was lost in the

when sodium carbonate is substituted under appropriate conditions for potassium carbonate. the relatively poor crystallising properties of

The probable mechanism of the ferrocyanide formation (Liehig, Annalen, 1841, 38, 20) is fusion by the action of potassinm carbonate on the organic nitrogen compounds present in the charred material (see Kraemer, B.P. 16529 of 1894, who describes the production of cyanide by heating carbazole to redness with causing potasb), together with iron sulphides produced from the sulphur containing bodies present in the organic material. Some iron sulphide may also be formed by reduction to sulphide of potassium sulphats present in the potash. For further details of the rôle of sulphur compounds in the fusion, the work of R. Hoffmann (Annales, 1860, 113, 81) may be consulted. The potassium cyanide and ferrous sulphide formed during the fusion react during the leaching to give potassium ferrocyanide.

6KCN+FeS=K4Fe(CN)8+K3S

2 From Coal Distillation Gases -The nitrogen z from Coal Distillation Cases—Ine nurgen content of bitummous coal varies widely, generally increasing with the geological age of the coal, but rarely exceeds 2%. The two types of coal which are subjected to distillation in practice are gas coal and coking cosl Gas coals generally contain from 1 to 2% of nitrogen, and nitrogen in various forms appears in all tha products (coke, tar, gas liquor and gas) of distillation (Drehschmidt, J. fur Gasbeleuchtung, 1904, 677; J. McLeod, J S C I 1907, 26, 137). The distribution of the nitrogen between these products varies considerably, being influenced by the nature of the coal, its moisture content and the conditions under which the distillation is carried out. A general indication of the nitrogen distribution in the distillation products is given by the following collected data on the distillation of small samples of dry coal.

		% of total N of the coal in				
No	Coal	Coke	NH ₃	нси	Tar	As N ₃
1 2 3 4	Westphaha		12-14 15	2	1·5 	35 2 30 0 35 19 5

Foster, J of Gas Lighting, 1882, 1081.

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than those actually produced in large-scale working, a partial explanation heing probably to be found in the fact that the experiments were carried out in retorts of iron, a metal which favours the decomposition of HCN at the distillation temperatures. A study of the numerous results of Drehschmidt (J. für numcrous results of Drehschmidt Gasbeleuchtung, 1904, 47, 677; see also Bertelsmann, "Technologie der Cyanverbindungen," Munich, 1906, p. 161) with English, West-phalian and Silesian coals does not disclose any connection between the nitrogen content of the coal and the distillation yield of HCN. On the other hand, as the water content of the coal (and the quantity of gas liquor obtained from it) increases, the yield of HCN decreases. This is due to the decomposition of HCN by steam at the distillation temperature into ammonia and carbon monoxide,

HCN+H2O=NH3+CO

(Carpenter and Linder, J.S.C.I. 1905, 24, 63). Thus the effect of the presence of 12% of moisture in a coking coal is to reduce by approximately one-third the amount of HCN present in coke oven gas as compared with town's gas

made from dry coal.

As regards the effect of distillation conditions, increasing temperatures cause a transfer of nitrogen from the coke to the more volatile products, leading to an increase in the yields of ammonia, HCN and gaseous Ammonia may be regarded as a primary decomposition product of the coal substance, and HCN as a secondary product formed by the interaction of ammonia with carhon:

$NH_3+C \rightleftharpoons HCN+H_2$.

This reaction is endothermic in the direction of HCN-formation, which is therefore favoured by higher temperatures (see Hydrocyanic Acid, p. 499). The following table shows the variation with temperature of the yields of ammonia and HCN obtained by O. Simmersbach (J.S.C.I. 1915, 34, 895) with a Silesian coal containing 1.396% of nitrogen:

Temp. (°C.) 600 800 1,000 1,200 % of total nitrogen---7.81 21.28 23.15 22.84 as NH₂ as HCN 0.250.871.231.42

According to Bertelsmann ("Technologie der Cyanverbindungen," p. 168) the size and nature of the carbonisation chamber influence the production of HCN, the large and porous chambers of coke ovens being disadvantageous by comparison with the smaller, relatively gas-

tight retorts of the gas works.

The gas leaving the distillation retorts contains ammonia, hydrogen sulphide, hydroeyanic acid (2-2.5 g. per cu. m.) and tar, and is first passed through coolers and scrubbers, the primary purpose of which is the removal of the ammonia and tar. In practice, however, considerable quantities of hydrogen sulphide and ammonium thiocyanate, and is ultimately unwanted thiocyanate at a later stage, but is

The yields of HCN quoted above are lower | discharged in the effluent from the ammonia stills; this effluent is sometimes worked up for thioeyanates, chiefly in England (Bertelsmann, op. cit. p. 180) via the cuprous and harium salts (see Thiocyanates, p. 508). The gas then passes to the iron oxide purifiers for complete removal of hydrogen sulphide, when 50-70% of the HCN is simultaneously taken up. This cyanide can be recovered by suitably treating the spent mass as described below (Dry Process). A more recent method of cyanide recovery from coal gas, giving greater efficiency, is the use of special cyanide washers, generally before the ammonia scrubbers (Wet Process).

(a) Dry Process.—The gas purification mass consists either of precipitated ferric hydroxide mixed with sawdust to increase the porosity, or of natural iron ores. The first technically successful mass was introduced by Laming (B.P. 11944 of 1847), and was made by mixing ferrous sulphate with slaked lime and sawdust and oxidising the damp mass in air. The use of natural ores was introduced by Howitz in 1870. The mass absorbs hydrogen sulphide from the gas, forming ferrous and ferric sulphides and elementary sulphur, and takes up HCN to form Prussian hlue and ammonium thiocyanate. When saturated with respect to hydrogen sulphide, the mass is regenerated by dampening and exposing to air, the ferrous and ferric sulphides heing oxidised to ferric hydroxido and sulphur. With successive regenerations the sulphur and cyanide contents of the mass increase until it is no longer effective as a

purifying agent.

The air-dried "spent oxide" is a dirty greenish, friahle substance smelling strongly of ammonia and organic sulphur compounds, and generally containing up to 50% of free sulphur and 10-14% of Prussian blue, except that the hlue content does not riso ahove 4% when a cyanide washer has heen used. This latter material is not considered attractive as a source of ferrocyanide, the minimum content at which spent oxide is salcable for ferrocyanide recovery heing about 7%. Ammonium thiocyanate may be present in quantities varying from traces up to 10%. The ferrocyanide in spent oxide is formed from the ferrous hydroxide and ammonium cyanide, and is then converted into Prussian hlue by interaction with the ferric iron. Ammonium thiocyanate is formed from hydrocyanie acid and ammonium polysulphide, the latter obviously arising from the presence of ammonia, hydrogen sulphide, and sulphur in the purifier; the formation of thiocyanate is favoured by raising the temperature, as occurs if the mass is regenerated quickly.

The different methods adopted for the treatment of the spent oxide are all variants of the following general procedure. The ammonium salts (chiefly sulphate and thiocyanate) are removed from the screened mass by systematic leaching with cold water for 14-24 hours, and the residue is dried. At this stage the elementary sulphur may be removed by up to one-third of the hydrocyanic acid (depending on the amount of water used in the scrubbers) bisulphide. This step is technically advanare also removed. This HCN appears as tageous, since it minimises the formation of

economically unsound owing to the fact that the and the recovery of ferrocyanide from "cyanide sulphur recovered is contaminated with tar mind" is considerably easier than from spent which can only be removed with great difficulty. The dry mass is then mixed in thin layers with the calculated amount of powdered slaked lime greater with gas which has already passed to decompose the Prussian blue content, the mixture warmed to drive off the last retained traces of ammonia, and then leached with water. An impure solution of calcium ferrocyanido is produced, from which the sodium or potassium salt may ho obtained by decomposition with the corresponding earbonate, filtering off the precipitated calcium carbonate and crystallising the solution. On the other hand, a Prussian bluo may be directly precipitated from the impure solution by acidifying with hydrochloric acid, removing the precipitate of sulphur thus formed, and adding ferrous or ferric salt solution, and the precipitated blue converted into potassium ferrocyanide by hoiling with caustic potash and evaporating the solution; this lengthy and expensive process is now used only for ferrocyanide recovery from

process to be described below

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Two improved methods of working up the impure calcium ferrocyanide solution were described by Kunbeim and Zimmermann (B P. 3342 of 1883), and are now generally used In the first, the sparingly soluble double salt K, CaFe(CN), is separated by the addition to the bot, concentrated solution (at 80°C) of the calculated amount of potassium chloride with constant stirring. The solid is filtered off and washed with a limited quantity of water, the filtrate and washings are worked up for potas-sium ferrogranide til Prussian hise by the method described above. The crystalline double salt is treated with the calculated amount of potassium carbonate solution at the boiling point, when calcium carbonate separates The elear potassium ferrocyanide solution is left to crystallise for 14 days, and gives directly a pure product. In addition to the purity of the product, this process has the advantage that one-balf of the calcium replacement is achieved with the cheaper potassium chloride instead of with the cheaper potassium chloride mistead of return protessium carbonate. The second improved of for method makes use of the spanning solubility of the plantaculum amnonia surface and in in this sea, the sufficient amnonia surface and present animonia surface neutralising with hydrochlorie acid. This salt is then decomposed with limo to give a pure calcium ferrocyanido solution, which is filtered sulphate solution: the ferrous sulphide sub-

oxide, It is also stated that the working efficiency of oxido hoves for H2S removal is through a cyanide washer, and that the spent oxide produced from such a gas is more suitable for sulphume acid manufacture (Drory, J. fur

Gasbeleuchtung, 1903, 46, 143). The first practical absorbent to he proposed was a suspension of ferrous hydroxide in an alkali carbonate solution (Knublauch, BP. 15164 of 1887, Ber. 1888, 21, 208), but the process did not meet with commercial success. Fouls (B.P. 9474 of 1892) substituted for ferrous hydroxido in the slurry a partially oxidised basic ferrous carbonate, the reagent being made by precipitating a ferrous salt solution with sodium carhonate, washing the precipitate and suspending it in sodium carbonate solution. This washing medium has achieved commercial success when used in a rotary washer designed to handle sturries (Foubs and Holmes, B.P. 1516s of 1895). When the cyanido washer is placed after the ammonia scrubber, the absorbed cyanids mother liquors and wash waters obtained in the appears in the form of soluble sodium ferrocyanide, which is recovered by evaporating the solution after filtering off the suspended solids A modified method of operation, practised at the Hague gas works, was described by Rutten (J. of Gas Lighting, 1902, 80, 875) and by Jorsseen and Rutten (J. fur Gasbeleuchtung, 1903, 48, 716), the crude gas, after tar sparation, is passed through the cyanide washer before going to the ammonia scrubher, and a 20% increase in the cyanide recovery results, both soluble and insoluble ferrocyanides being formed. The saturated liquor from the washer, contaming Prussian blue, potassium and ammonium ferrous ferrocyanides in suspension, and ammonia and potassium ferrocyanides in solution, is filter pressed, and the filtered solution distilled to separate ammonia and then evaporated in tacuo to erystallisation. The press-cake is boiled with caustic potash and the ammonia collected, the iron oxide being returned to the cyanids washer and the solution of ferrocyanido proceeding to the evaporation

The ammonia present in the gas may itself be used as the alkalı and the crude gas be simply washed with a ferrous sulphate solution. In this case, ammonia and hydrogen sulphids present in the gas give ammonium sulphide, so that the wash liquor is quickly converted into a euspension of ferrous sulplude in ammonium calcium ferrocyanido solution, which is nueved supprite solution; the secretary supprise solution and treated by the KCl+K_CO_p process sequently racets with animonium cyanide in already described. The process or pure.

(b) Flet Process—The wet process of cyanide removal first came into successful practical R.P. 2083 of 1896; Schroder, B.P. 19849 of operation in 1895, and is now an important 1898; Telehmann, BP. 12485 of 1899) used source of ferrocyanides. It has several advanding ferrous salt solutions so that ammonis was source of ferrocyanines. As an several available that tages over the dry process. The absorption always present in excess, with the object of HCN in purifier hoxes is an unroluntary obtaining ferrocyanide wholly in the soluble adjunct to the main purpose of sulphar removal, forms. Buch, on the other hand (BL. 2012). and the removal of HCN is not complete.

The cyanido scrubber, on the other hand, of 1893), almost completely avoids the formation of soluble salts by using a saturated ferous proudes a controlled method of cyanide removal subhate colution, maintaining ferrous groups. which works at an efficiency of nearly 100%; in permanent excess in the solution and pro-

ducing ferrocyanide in the insoluble form. This process is widely used, and is said to absorh 98% of the hydrocyanic acid in the gas to give a mud containing the equivalent of 18-20% of K₄Fe(CN)₆ (Bueh, J. für Gasheleuchtung, 1900, 43, 747). The mud is treated with some ferrous sulphate solution to convert small quantities of soluble ammonium ferroeyanide into insoluble ferrocyanide, and then with steam to remove ammonia. It is then dried in filter presses and the cake, containing the equivalent of 40% of K4Fe(CN)6, is eonverted into calcium ferrocyanide solution hy boiling with milk of lime. The clarified solution is then converted into potassium ferrocyanide by the method of Kunheim and Zimmermann already described.

The mechanism of ferrocyanide formation in the wet absorption of cyanide from coal gas, and an explanation of the effect of the variation of the conditions on the form in which the product appears, have been given by Feld (J. für Gasheleuchtung, 1904, 47, 132). He considers that the basic reaction is between ammonium cyanide and ferrous sulphide, formed in the way previously explained. If ferrous sulphide is present in excess, insoluble ferrous ferro-cyanide is the main product; under more balanced conditions, insoluble ammonium ferrous ferrocyanide is formed in increasing quantities, and when there is a deficiency of ferrous sulphide, the soluble ammonium ferro-

cyanide is produced.
3. From Thiocyanates.—Many processes have been described for the conversion of thio-cyanates, which are formed in various methods of eyanide synthesis and in gas purification, into the more valuable ferrocyanides. processes consist essentially in a desulphurisation with metallic iron, and may he carried out with the molten salts or their solutions. Alexander (J. für Gasbeleuchtung, 1878, 21, 20) described a method in which ammonium thiocyanate, potassium carbonate, carhon, and iron filings are rubhed to a paste with oil and heated in a closed iron vessel to red heat. The reaction product is leached with an aqueous suspension of freshly precipitated ferrous hydroxide, and the filtrate evaporated to crystallisation. Tscherniac and Gunzburg (B.P. 1261 of 1881, see also 1148 of 1878 and 1359 of 1879) and Hetherington and Muspratt (B.P. 5830 of 1894) used a 20-30% excess of reduced iron and molten potassium thiocyanate which had heen dried at 300°C., the reaction temperature being 445°C. Crowther, Rossiter, Hood and Albright (B.P. 8305 of 1894) recommend drying the thiocyanatc in a stream of nitrogen or earhon dioxide. In laboratory experiments on Hetherington's method, Conroy (J.S.C.I. 1896, 15, 12) obtained a 77% conversion of thiocyanate to ferrocyanide. The mechanism of the reaction is analogous to that of ferrocyanide formation from nitrogenous organic matter, viz. cyanide and ferrous sulphide are formed during the fusion, and react to give ferrocyanide during the leaching.

The desulphurisation may he carried out in

sium thiocyanate with twice the theoretical weight of iron filings necessary for sulphide formation and twice the theoretical weight of ferrous hydroxide necessary for ferrocyanide formation, at 110-120°C. under pressure. In 12 hours a conversion of 80% is obtained. Conroy (J.S.C.I. 1898, 17, 98) used a mixture of calcium thiocyanate and ferrous chloride and iron in water at 140-150°C. to increase the rate of reaction:

$$\begin{array}{l} \operatorname{Ca(CNS)_2} + \operatorname{2Fe} + \operatorname{FeCl_2} \\ = \operatorname{CaCl_2} + \operatorname{Fe(CN)_2} + \operatorname{2FeS} \end{array}$$

The mud is treated with hydrochloric acid to dissolve ferrous sulphide, and the ferrous ferroeyanide hoiled with alkali to form soluble ferro-

Copper (Bower, B.P. 361 of 1896) or nitric acid may he used as desulphurising agents (see

Hydrocyanic Acid, p. 492).

4. From Cyanides .- The fact that the conversion of synthetic cyanides into ferrocyanides is becoming a technical process displacing the recovery of by-product ferrocyanide at gas works, especially in the United States, indicates the change in the relative economy of cyanides and ferroeyanides which was caused by the introduction of cyanide synthesis processes from ammonia and from nitrogen. The technical conversion of cyanide to ferrocyanide is brought about by the action of cyanide solution on iron, iron sulphide or ferrous salt. The chief source of cyanide for this purpose is the crude calcium cyanide (analysing 45% Ca(CN)2, 32% NaCl) obtained on fusing technical calcium cyanamide with sodium chloride (see Alkaline Earth Cyanides, p. 488). This is dissolved in water and treated with the calculated amount of ferrous sulphate to satisfy the requirements of the reaction equation

$$3Ca(CN)_2 + FeSO_4 = CaSO_4 + Ca_2Fe(CN)_6$$

and with slightly less than the amount of sodium carbonate equivalent to the ferrocyanide produced:

$$Ca_{2}Fe(CN)_{6}+2Na_{2}CO_{3}$$

$$=Na_{4}Fe(CN)_{6}+2CaCO_{3}.$$

The sodium ferrocyanide in solution is separated from the sodium chloride by crystallisation (Washhurn, U.S.P. 1398453, 1921; Rugh, U.S.P. 1650390, 1927).

The American Cyanamid Co. (U.S.P. 1667839, 1928) proposes to make the slightly soluble calcium potassium ferrocyanide hy a similar reaction after addition of potassium chloride. The straightforward interaction of sodium eyanide (the commonest synthetic cyanide) with ferrous sulphate presents some difficulty in the separation of the two soluble products, sodium ferrocyanide and sodium sulphate. According to Dominik (see Chem. Zentr. 1923, IV, 494), the solubility relations are such that sodium sulphate is retained in solution on evaporation above 32°C. For the conversion of sodium ferrocyanide into the potassium salt, which is sometimes required, it is necessary to solution hy tho method of Sternberg (G.P. add to the hot solution a considerable excess of 32892 of 1882), hy heating a solution of potasadd to the hot solution a considerable excess of

212698, 1909). Instead of using calcium cyanide made from cyananide, the Strontian- und Potasche-Fabrik (G.P. 330194, 1920) describe a method

of converting eyanamide directly into ferro-cyanide by heating it to redness with potassium carbonate and iron shavings in the absence of air

PROPERTIES OF THE FERROCYANIDES.—In this section the principal characteristics are described of those selts which possess technical interest. For further information, "Cyanogen Compounds" by H. E. Williams (1915) or an "Inorganic Chemistry" eg. Mellor's should be consulted.

Ammonium ferrocyanide is prepared in aqueous solution from the free acid and ammonia, from barium ferrocyanide and ammonium sulphate, or from lead ferrocyanido and ammonium carbonate. The solid salt, (NH4)4Fe(CN)4.3H2O, is obtained by evapo rating the solution in secue (Briggs, J.CS 1911, 99, 1019), or by precipitation with alcohol (Cumaning and Good, J.C.S. 1926, 129, 1924). The salt crystallises with difficulty in very pale yellow plates, isomorphous with the potassium salt (Bunsen, Pogg Ann 1835, 36. 404) When (NH4)4Fe(CN)6 is heated in 1 acue, decomposition begins at 110°C., NH4CN being evolved (Mittageh, Kuss and Emert, Z anorg Chem 1928, 179, 193); at 320°C, "Fe(CN), " remains behind, which begins to evolve nitrogen at 430°C; and at 700°C a mixture of iron, iron carbide and nitride remains. When the heating is carried out in an atmosphere of hydrogen, methane and ammonia are formed and the residue is pure iron. The aqueous solution decomposes on boiling to give ammonium cyanide and an insoluble residue possibly consisting of partly exidised ammonium ferrous ferro yanide

From solutions containing much ammonium chloride a double salt,

(NH₄)₄Fe(CN)₆ 2NH₄Cl₅3H₂O₄

crystallises readily in well-formed orange yellow rhombohedia, which were formerly mistaken for the ammonium salt (Bunsen, le).

Barlum ferrocyanide, Ba, Fe(CN), 6H,O, is precipitated on moving strong solutions of barium chloride and sodium ferroes anide (Ewan and Namer, JSCI 1913, 32, 467) as a light yellow, crystalline powder. The prepara-tion of pure barum salt from K,Fe(CN)₈ as difficult, as the product generally contains potassium; Walker (J. Amer Chem. Soc. 695, 17, 927) avoided the difficulty by first making the dimethylandine salt and decomposing this with burnini hydroxide, extracting the liberated dimethylandine with ether. The dried salt, heated in tacuo or in nitrogen, begins to decompose at about 500°C., giving nitrogen, carbon, iron, and barium eyanide and cyanamide When oxygen is completely excloded, the molecular ratio of cyanamido to evanide in the product is 2:1 (Ewan and Napier, Ic.) The solubility of the salt in water, expressed as g. of anhydrous salt in 100 g. of water, is 0 34 at 15 5°C. and 1 01 at 100°C. More than 5 molecules of water of crystallisation are rela and Kellerholf, J. pr. Chem. 1912, [11], 86, 82)

(Conroy, J.S.C.I. 1898, 17, 98; Petri, G.P. | tively easily removed by heating to 80°C, for a few hours, but the remainder is not completely given up, even at 180°C (Ewan and Napier, Ic). Barium potassium ferrocyanide.

is precipitated as a pale yellow, crystalline powder on mixing equivalent quantities of dilute solutions of barium chloride and potassium ferrocyanide. The crystals effloresce readily, a property which accounts for the uncertainty as to the amount of water of crystallisation solubility in water (g of anhydrous salt in 100 g of water) is 2 1 at 15 5°C and 6 08 at 100°C. Calcium ferrocyanide,

Ca,Fe(CN),11H,O.

is made by treating Prussian blue with milk of lime, neutralising hydroferrocyanic acid with calcium earbonate, or from potassium ferro cyanide ma the dimothylanibne salt (see Barium Perrocyanide, above) The crystals were formerly believed to contain 12H.O. but more recent work (Berkeley, Hartley and Button, Phil Trans. 1909, 209, (A), 177; Cumming, JCS 1921, 125, 240, Farrow, bid. 193, 123, 49, 1927, 1153) favours 11H,O. The salt is extremely soluble in water (cf. the barrum salt), giving viscous solutions at high concentrations. The crystals efflusees and lose concentrations. The crystals eministee and lose all but i molecule of water at 40°C, Cumming (t c) detected some hydrolysis to give a compound CapFeg(CN)₁₁OH on heating at 100°C in a current of oxygen-free gas, At 500°C in the absence of oxygen, Pincass (Chem. 2tg. 1922, 48, 681) observed a decomposition of the control of the composition of the control of the co position to give calcium cyanamide.

Ca.Fe(CN),=2CaCN.+Fe+N.+4C Calcium ammonium ferrocyanide.

Ca(NH₄)_eFe(CN)_a, may be prepared from calcium chloride and ammonium ferrocyanide solutions, and forms a wlute, minutely crystalline powder, containing no water of crystallisation, which becomes coloured on exposure to light. 100 g of water at 15-17°C dissolva 0 258 g of the salt [J. Campbell Rosso, J.C.S. 1907, 91, 1826), The insolubility of the salt is used in separating

ferrocyanide from gas purification mass. Calcium potassium ferrocyanide,

CaKaFe(CN)...

is made from calcium chloride and potassium ferrocyanide solutions, forming a white powder of amail quadratic prisms which are anhydrous and discolour slightly on exposure to light. At 15-17°C the adjubility is 0 41 g, per 100 g of water (J Campbell Brown, Ic).

can be prepared either from cupric salt and ferrocyanide, or from cuprous salt and ferrieyanide, for the ionic oxidation reduction equilibrium.

is almost completely to the left (Müller, Wegelin

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addition of a solution of calcium ferrocyanide or bydroferrocyanic acid to an excess of cupric chloride, and the product is a red-brown, substance containing amorphous amounts of water. It is very slightly soluble in water and dilute acids, but can be brought into colloidal solution by means of peptising agents, particularly through the agency of ferrocyanide ions adsorbed from solutions. The brown precipitate formed when cupric salts react with ferrocyanide (Hatcbett's brown, potassium formerly used as a pigment) always contains potassium, even when cupric copper is present in cxcess, and this may be explained either by adsorption of K_4 Fe(CN)₆ by the solid Cu₂Fe(CN)₆ (Luckow, Chem.-Ztg. 1892, 16, 164, 836, 1428; Duclaux, Compt. rend. 1904, 138, 144, 571) or by the formation of a double salt (c.g. CuK₂[Fe(CN)₆],2H₂O, Reihlen and Zimmermann, Annalen, 1927, 451, 75). Colloidal membranes of copper ferrocyanides have been used extensively in the study of osmosis, and copper ferrocyanide finds application in photo-graphy for toning: the black deposit of silver in a hromide print is converted into a red one, consisting of copper ferrocyanide, by immersion in a solution of copper sulphate, potassium ferricyanide, and potassium citrate (Ferguson, Phot. J. 1900, 133).

Reihlen and Zimmermann (Annalen, 1927, 451, 75). regard cupric ferrocyanide as the copper salt of a complex anion containing hoth copper and iron; for on hoiling with milk of lime it gives brown crystals of Ca[CuFe(CN)6], and with dilute ammonia it gives the salt [Cu(NH₃)₄][CuFe(CN)₆]H₂O; such reactions indicate that one-half of the copper in cupric

ferrocyanide is in the complex ion.

Cuprous ferrocyanide is not known in the simple form, but the existence of the double salts KCu₂Fe(CN)₆ and K₂Cu₂Fe(CN)₆ has heen indicated by Müller, Wegelin and Kellerhoff (J. pr. Chem. 1912, [ii], 86, 82), using the method of indirect analysis described later. Messner (Z. anorg. Chem. 1895, 8, 368) describes the preparation of $K_2Cu_2Fe(CN)_6$ by the action of cuprous cyanide or cuprous chloride, in the presence of potassium sulpbite to prevent oxidation, on potassium ferrocyanide solution, the mixture being quickly cooled to obtain the double salt in the form of colourless plates. It appears likely that, in the above double salts, the copper is part of the complex ion (Reihlen and Zimmermann, Annalen, 1927, 451, 75). The cuprous ferrocyanide compounds differ from other ferrocyanides in evolving the whole of their cyanogen as bydrocyanic acid on boiling with dilute sulphurie acid (see Dewrance and Williams, B.P. 28074 of 1908).

Hydroferrocyanic acid, H4Fe(CN)6, may he prepared by the decomposition of a suspension of the lead salt with sulphuric acid and evaporation of the filtrate, or by treating a concentrated solution of potassium ferrocyanide with excess of concentrated hydrochloric acid, prefcrably with addition of ether (see, for example, Mittasch and Kuss, Z. Elektrochem. 1928, 34, 159). The acid crystallises in colourless scales which are

The usual method of preparation is by the dried over sulphuric acid. It may be purified by dissolving in alcohol and precipitating with ether; it obstinately retains ether, with which it forms a compound (see, for example, Cumming and Brown, J.S.C.I. 1928, 47, 84T), and must be beated to 100°C. in a stream of dry hydrogen to free it completely (Browning, J.C.S. 1900, 77, 1234). Heated in the absence of oxygen, the acid begins to decompose at 150°C. into HCN and "Fe(CN)2." The solubility of the acid in water at 14°C. is 13 g. per 100 g. of solution. The solution is readily oxidised by air at higher temperatures, forming a blue substance which is assumed to be Prussian hlue; this reaction is used in calico printing for the production of hlue patterns.

Ferrous ammonium ferrocyanide,

$Fe(NH_4)_2[Fe(CN)_6],$

formed when ammonium ferrocyanide solution is exactly precipitated by a ferrous salt in the presence of an excess of an ammonium salt, oxygen being excluded. It is also formed when ammonium ferrocyanide is distilled with an acid in the absence of air.

Ferrous potassium ferrocyanide,

K₂Fe[Fe(CN)₆],

(Everitt's salt) is produced by heating saturated potassium ferrocyanide solution for 40 hours at 90°C. with an equal volume of 20% sulphuric acid, washing and drying the residue with exclusion of air (Hofmann, Heine and Höchtlen, Annalcn, 1904, 337, 1). The pale yellow product is not oxidised hy air, and is therefore different from the isomeric salt obtained when solutions of potassium ferrocyanide and ferrous chloride are mixed (Hofmann, l.c.). For this reason Reihlen and Zimmermann (Annalen, 1927, 451, 75) formulate it as $K_2[Fe_2^{II}(CN)_6]$.

Iron cyanide blue compounds.-The hlue compounds of iron and cyanogen may he prepared hy the interaction of ferric salts with ferrocyanides, or of ferrous salts with ferricyanides, in neutral or acid solution. The constitution of these compounds is discussed in greater detail on p. 473, but it may be stated here that they are believed to he ferrie ferroeyanides. Müller bas shown that the composition of the unwashed precipitate initially formed depends on the relative quantities of reagents employed (Chem.-Ztg. 1914, 38, 281, 328). Starting from potassium ferrocyanido and ferric chloride, Prussian blue is precipitated when the molar ratio K_4 Fe(CN)₆/FeCl₃ is less than 0.75; for values between 0.75 and 0.92 the precipitate contains increasing quantities of KFe^{III}(Fe^{II}(CN)₆], and when the ratio exceeds 0.92 the substance K₂Fe^{II}[Fe^{II}(CN)₆] is also present. Starting from potassium ferricyanide and ferrous chloride, a precipitate of the composition KFe₂^{II}Fe₅^{III}[Fe^{II}(CN)₆]₅ is obtained when the molar ratio K3Fe(CN)6/FeCl2 is less than 0.714; between 0.714 and 0.75 a substance KFe^{II}Fe₃^{III}[Fe^{II}(CN)₆]₃ is formed, which Müller regards as ordinary *Turnbull's blue*; from 0.75 to 0.9 increasing quantities of KFe^{III}[Fe^{II}(CN)₆] are present, and finally, when the ratio exceeds 0.9 ordinary Prussian washed with water saturated with ether, and blue, Fe4III[FeII(CN)6]3, appears.

Blue compounds are also obtained by the hydroferrocyanic acid, H4[FeII(CN)4], or as oxidation of ferrous ferrocyanide, and by the reduction of ferric ferricyanide. All the blue compounds are amorphous substances containing indefinito quantities of water, some of which (see Constitution of the Blue Compound, is so tenaciously retained as to be regarded as p. 473), and in the following section the formula water of constitution. Their colour depends on the both systems are quoted. the composition, water content, and particle size, and varies from pure blue to deep violet blue, a coppery metallic surface reflection being sometimes also present They do not form true solutions in water. The so called "aoluble blucs" give deep blue colloidal solutions with pure water, from which they can be precipitated by the methods normally effective with colloidal solutions. "Insoluble blues" are not peptised in pure water, but only in oxalic acid solution, and sometimes in acid or neutral ovalates and

The blue compounds are quite stable towards dilute acids Concentrated hydrochloric and sulphuric acids dissolve them with decomposition, giving ferrous or ferrie salt and hydroferro. or hydroferri cyanie acid, the blue may be re formed on diluting the solution. They are coagulated by alcohol Alkalis, or salts with an alkaline reaction, decompose them to give iron hydroxide and ferrocyanide, the rate of decom position varying with the different compounds and providing a method of distinguishing them Colloidal solutions of Prussian blue (0 2 to 2% are negatively charged, the blue being coagulated by kations, especially hydrogen ion

The characterisation of different blues by analysis is difficult. Simple analytical methods are useless because the collindal nature of the blues gives them the property of strongly adsorbing dissolved salts from solution, and renders the usual methods of purification generally impracticable. In addition, some blue compounds, which contain ferrous iron outside the complex radicle, are easily oxidised by stmospheric oxygen, so that changes of com position may occur during the manipulation of the original product, unless special pro-cautions are taken. For this reason, the indirect method of analysis, by which the composition of a precipitate is determined by mixing known quantities of the reagents and estimating the excess left in the supervisiont hound, has been used by Müller and his school (Muller and Stamsch, J. pr. Chem. 1909, [n], 79, 81; 1909, [n], 80, 153; Muller, Wegelm, Treadwell and Diefentbaler, ibid 1911, [11], 84, 353; Muller, ibid. 1914, [11], 90, 119) and by Woringer (1bid. 1913, [11], 87, 51), although the objection still holds that the precipitate is not a pure

aubstance on account of adsorption. A description will now be given of the more important types of blue compounds, the data referring in each case to the solids obtained after exhaustive drying of the original precipitates with phosphorus pentoxide. Our know-ledge of these compounds is largely based on the work of Hofmann and his collaborators (Hofmann, Heme and Hochtlen, Annales, 1904, 337, 1; Hofmann and Resenscheck, shid. 1905, 340, 207; 1905, 342, 364; Hofmann, Arnold, and by oxidising ferrous ferrocyanide in the presence for the first state of the sta

salts of a hydro-ferro ferricyanic acid, H[FeHFeH(CN),]

p. 473), and in the following section the formula for both systems are quoted. Soluble Prussian blue.

> KFefffFeff(CN), aq or KiFeliFelli(CN) alag.

is formed when cold neutral solutions of potas sum ferrocyanide and ferric chloride are mixed. the ferrocyanide being in excess; the precipitate is washed with aqueous alcohol to free it from electrolyte, and dried in tacue over sulphune acid or phosphorus pentoxide The same substance is also formed by adding cold neutral ferrous salt to an excess of potassium ferri examide solution, and similarly treating the precipitate (Skraup, Annalen, 1677, 186, 371, Hofmann, Heine and Hochtlen, Annalen, 1904, 237, 1) The water content of the dried substance is variously reported as up to 4H.O. of which 2 molecules cannot be removed without decomposition, and are therefore regarded as constitutional. The substance is soluble in water, giving a colloidal solution which is coagulated by adding strong electrolytes or acrds, and by alcohol. It is also soluble in ovalic acid solution

Substances of similar composition and but slightly different properties are obtained by treating ferrous sulphato solution with potassium ferrocyanide, and oxidising the white ferrous ferrocyanide produced, either with atmosphere oxygen or with hydrogen peroxide. When this is done in neutral solution, the product,

KFeIII[FeII(CN),],H,O,

is soluble in water, but not in ovalic acid solution and is immediately decomposed by ammonia solution (Hofmann) When the preparation is carried out in the presence of 1% of sulphune acid, on the other hand, it yields an isomeric blue compound which is soluble in water and in ovalie acid solution, but is more stable towards ammonia solution, requiring 6-7 minutes for the precipitation of ferric hydroxide,

A whalse blue containing ammonium instead of potassium (Monthier's blue) can be obtained by oxidising ferrous ammonium ferrory anide untb air or hydrogen peroxide.

Insoluble Prusslan blue,

Fe, III (FeII (CN), lag or Felit(FeliFelii(CN),]aq,

can be made by decomposing cold neutral or alightly acid potassium ferrocyanide solution with excess of ferric salt, washing the pre cipitate with dilute hydrochloric acid and drying it in vacuo. It is also produced from soluble Prussian blue by digesting with ferric chloride : colution

3K[FeHFeH (CN),]+FeCl, =Fetti FettFett (CN) 3+3KCI;

water content of the product varies considerably according to the treatment during preparation, but after long drying in vacuo over phosphorus pentoxide reaches a lower limit of $10-11\,H_2O$ (Hofmann, Heine and Hochtlen, Annalcn, 1904, 337, 1). Further dehydration by heating above 150°C. is accompanied by decomposition. Insoluble Prussian blue does not give colloidal solutions with pure water, but only with oxalic acid, oxalate and tartrate solutions.

Oxidising agents such as chlorine and nitric acid frequently give rise to green solutions and green precipitates ("Prussian green"), which are believed to be complex mixtures intermediate in composition between ferric ferrocyanide and ferric ferricyanide.

Insoluble Turnbull's blue,

 $\begin{array}{l} {\sf Fe^{II}Fe_2^{III}[Fe^{II}(CN)_6]_2}{\rm aq.~or} \\ {\sf Fe^{II}[Fe^{II}Fe^{III}(CN)_6]_2}{\rm aq.,} \end{array}$

is produced by precipitating a solution of potassium ferricyanide with excess of ferrous salt, or by treating soluble Prussian blue with ferrous salt:

$$\begin{split} 2\mathsf{K} [\mathsf{F} \mathrm{e}^{\mathrm{II}} \mathsf{F} \mathrm{e}^{\mathrm{III}} (\mathsf{C} \mathsf{N})_6] + \mathsf{F} \mathrm{e} \mathsf{C} \mathrm{I}_2 \\ = & \mathsf{F} \mathrm{e}^{\mathrm{II}} [\mathsf{F} \mathrm{e}^{\mathrm{II}} \mathsf{F} \mathrm{e}^{\mathrm{III}} (\mathsf{C} \mathsf{N})_6]_2 + 2\mathsf{KCI}. \end{split}$$

It is formed by the reduction of insoluble Prussian blue with ferrous ion or sulpbite (Müller, J. pr. Chem. 1914, [ii], 90, 119). The water content of Turnbull's blue is variously reported. The blue is colloidally soluble in oxalic acid solution, but not in water. Moderate oxidation, or reaction with ferrie salts, gives insoluble Prussian blue:

 $\begin{array}{l} 3 Fe^{II}[Fe^{II}Fe^{III}(CN)_6]_2 + 2 FeCl_3 \\ = 2 Fe^{III}[Fe^{II}Fe^{III}(CN)_6]_2 + 3 FeCl_2 \end{array}$

Williamson's blue or violet,

KFe^{III}[Fe^{II}(CN)₆],H₂O,

is obtained by the oxidation of Everitt's salt, K₂Fe^{II}[Fe^{II}(CN)₆] with warm dilute nitric acid, or with an excess of hydrogen peroxide in the presence of dilute sulpburic acid (Hofmann, Heine and Höchtlen, Annalen, 1904, 337, 1). It is a violet-blue substance, not peptisable in water or in oxalic acid solution, giving only a greenish-blue turbid suspension with water. It is relatively unreactive, giving ferric hydroxide with 4% ammonia solution, for instance, only after some hours. It is thus entirely different from the isomeric blues obtained by oxidising ferrous forrocyanide (see Soluble Prussian Blue, above). It is a general rule that the precipitation of blues from hot acid solutions tends to produce substances of a more stable type, approaching Williamson's blue in properties.

Commercial blues are usually mixtures of the compounds already described, and appear in commerce under such names as Paris blue bronze blue and steel blue (mostly containing potassium, and of superior properties), Prussian blue and mineral blue (less valuable products consisting of mixtures), and gas blue or Monthier's blue (containing ammonia). Of the three [ii], 89, 51) was indeed able to show that ferri-

iron outside the complex radicle (see below) salts with ferricyanide (formation of Turnbull's (Müller, Chem.-Ztg. 1914, 38, 281, 328). The blue bas been discontinued, and the reaction of blue) bas been discontinued, and the reaction of ferric salts with ferrocyanide (formation of Prussian blue) is rarcly practised. The best blues are made by precipitating a ferrous salt with potassium ferrocyanide and then oxidising the precipitate (Gentelc, "Lehrbuch der Farbenfabrikation," 1906). Paris blue, the finest quality of commercial Prussian blue, may be made by mixing 45 kg. of FeSO₄,7H₂O (a slight excess), dissolved with exclusion of air in 250 kg. of water, with 50 kg. of K₄Fe(CN)₆ dissolved in the same quantity of water, and running the mixture into a further quantity of 250 kg. of water. The precipitate is washed and filtered off to be oxidised by suspending it in water, dilute sulphuric acid or ferrous sulphate solution and treating it with nitric acid of sp.gr. 1.23, ferric obloride solution (not in excess), cblorine, or potassium chlorate solution. precipitate of Prussian blue is then washed with water until free from sulphate or chloride, filtered and air-dried at 30-40°C. Chlorate is now said to be the preferred oxidising agent (Itzkovich and Shmul'yan, see Amer. Chem. Abstr. 1934, 28, 3997).

Paris blue contains potassium ferric ferrocyanide, and the presence of the potassium leads to an enhancement of the properties of the blue. If sodium is present instead (by making the preparation from sodium ferrocyanide) a less satisfactory colour is produced. Eibner and Gerstacker (J.S.C.I. 1912, 31, 1041) measured the potassium and water contents of a number of blues: steel blue and bronze blue eontained 12% K and 5.5% H₂O, Paris blue 9% K and 10.5% H₂O, commercial Prussian blue 1.5% K and 24% H₂O. The rise of water content as the potassium content falls is marked, and may be explained as an effect of the electro-lyte (potassium ion) in "salting out" the colloid and reducing the water absorption.

In commercial pigments, Paris blue is diluted with finely ground starch, heavy spar, gypsum, zinc white or burnt kaolin, and the mixture milled. Blues of very low quality are faced by rotating the dry cubes in a cask with a fine dust of pure Paris blue.

Constitution of the Blue Compounds.—It is certain that all forms of blue contain both FeII and FeIII, since ferrous ferrocyanide is white, and ferric ferricyanide is brown and soluble. Müller and Stanisch (J. pr. Chem. 1909, [ii], 79, 81) and Müller (Chem.-Ztg. 1914, 38, 281, 328) calculated from the known normal potentials of the ionic reactions:

$$Fe^{++} \rightleftharpoons Fe^{+++} + \Theta$$

 $Fe(CN)_6'''+ \ominus \rightleftharpoons Fe(CN)_6''''$

that the equilibrium constant of the reaction

$$Fe^{++}+Fe(CN)_6'''' \rightleftharpoons Fe^{++}+Fe(CN)_6''',$$

 $k(=[Fe^{++}][Fe(CN)_6'''']/Fe^{++}][Fe(CN)_6'''])$

is approximately 10⁻⁵, so that, in a solution containing all these ions, the equilibrium would be practically completely on tbe side of ferrous ferricyanide. P. Woringer (J. pr. Chem. 1913, methods of manufacture, the reaction of ferrous cyanide is formed when ferric chloride reacts

that a ferricyanide structure must be assigned. They apply the same principle to other insoluble to the blue compounds. It is not permissible, ferrocyanides such as the cupric and zinc salts. however, to deduce from solution equilibria the the Ferri in the blue complex being replaced in composition of a precipitate, which is controlled these cases by Cu^{II} and Zn^{II} respectively, and by solubility relationships. If, for example, the to double salts of ferrocyanide, such as Eventt's oy somonity relationships. If, for example, the ratio of the solubility products of ferric ferroganide ([Fe+++]Fe(CN],"") and ferrous ferricyanide ([Fe+++]Fe(CN],"") is less than $l(=10^{-4})$, then practically complete precipitations. tion of ferrie ferrocyanide could occur from a solution containing ferrous and ferricyanide ions, in spite of the fact that the ionic equilibrium is entirely in favour of the alternative compound.

It follows that only one of the two alternative compounds is stable, that mixtures of the two cannot exist in contact with solution, and that the composition of an equilibrium product must always be either ferrous ferricyanide or ferric ferrocyanide, independent of the starting msterials. The ratio of the two solubility products is unknown, for neither has been measured. On the basis of the Abegg-Bodlander theory of electro-affinity, however, we may pre-dict that the solubility of the compounds formed from the more highly charged ions Fe+++ and Fe(CN),"" of weaker electro-affinity will bo considerably lower than that of the compounds from the ions Fe++and Fe(CN),", of lower electrovalency and stronger electro-affinity, so that the blue compounds are probably ferro-

oyanides.

The chemical evidence must be interpreted with eaution. The conclusions of Hofmann and his collaborators (Annalen, 1904, 237, 1, 1907, 352, 54) that insoluble Prusman blue is ferrio ferrocyanide, because it is formed from ferrio ferricyanide and hydrogen peroxide, which reduces ferricyanide and not ferric son, and of Eibner and Gerstacker (Chem. Ztg. 1913, 37, 137, 138, 195) that the blue compound made by the sulphite reduction of ferms ferricyanide is ferrous ferricyanide, because sulphite reduces only ferme ion, are pasound because the alternative products might be formed from the original reduction products in each case by the operation The best reason for of Müller's equilibrium. regarding insoluble Prussian blue as ferrie ferrocyanide is that it is not oxidised by oxygen or by hydrogen pecoxide, so that its divalent iron is in the complex: its method of formation from ferrio salt and soluble Prussian blue suggests that the soluble blue is also a ferrocyanide: with introgen escaping. According to Hackspill and since Turnbull's blue is produced from and Pinck (Bull, Soc. chim. 1931, [17], 49, 54) aoluble Prussian blue and ferrous salt, it must also | considerable quantities of eyanogen are obtained

be a ferrocyanide. The conception of Prussian and other blues as giving the simple iron ferrocyanides is not entirely satisfactory, however, because it leaves unexplained their differing degrees of stability towards 100 g. of solution; alkalıs and oxidising agents, and does not account for their remarkably deep colours, which would not be expected of compounds formed by the simple combination of feebly-coloured ions (Hantzsch, Z. physikal. Chem. 1910, 72, 362). Reihlen and Zimmermann (Annalen, 1927, 451, 75; 1929, 475, 101) and Reihlen and von Kummer (Annslen, 1929, 469, 30) have put forward the view that the iron cyanide complex in the blue compounds contains both pletely decomposed by boiling with mercune

with an excess of ferrocyanide, and concluded | Fe^{II} and Fe^{III}, with which CN is ro ordinated. salt, K₂Fe[Fe¹¹(CN)₆], which they write as K₂[Fe¹²₂(CN)₆] This method of formulation simplifies considerably the constitutional problems of the iron cyanogen blue compounds. Only two alkah free compounds are possible: (a) Fem[FemFem(CN)al, in which one seventh of the total iron is reducible by sulphite. and which is readily identified with insoluble Prussian blue, and (b) FeII[FeIIFeIII(CN)] m which one-fifth of the total iron is oxidisa by hydrogen pecoxide, and which is Turnbull's hine; and soluble Prussian blue is the potassium salt of the ferro ferri complex

K[Fe^{TI}Fe^{TI}(CN),].

The interconversions of soluble and insoluble Prussian blues and Turnbull's blue are readily explained. Cambi and Clenci (Gazzetta, 1928. 53, 57) accept the theory of the complex con-taining both divalent and trivalent metals coordinated with cyanide for all coloured complex cyanides; for example, the bluisb-green ferrous manganicyanide, usually written

Fe,"[MnIII(CN),]2,

decomposed by alkalı into manganic is decomposed by sikal into manganic hydroxide, Mn(OH), and ferrocyande, whereas the colourless ferrous cobalticyande Fe₃¹²(Co¹¹(CN)₈)₂; gives ferrous hydroxide, Fe(OH)₈ and cobalticyande. Hence the rolour: less cobalt compound is correctly formulated above, but the coloured manganess compound should be written MnIII (FeIIMnII (CN),],

or as a similar more complex derivative, Potassium ferrocyanide, "yellow prussiate of potash," K,Fe(CN),3H,O, crystallises from aqueous colutions in large, mouoclinic tetragonal pyramids of sp gr. 186. The crystals are stable in air at the ordinary temperature, but begin to lose water at 60°C., being completely debydrated at 100°C. to give a white powder. On further heating in the absence of exygen, the powder decomposes, first mto Williamson's salt, KrFe[Fe(CN)], without gas formation, and then below red heat into a mixture of potassinm cyanide, carbon and iron. at 300-400°C. The salt is soluble in water, following saturated solutions (Farrow, J.C.S. 1926, 129, 49); the solubilities

are expressed as grams of anhydrous salt in Temp. °C. Solubility 249 349 498 33 I3 23 95 28 OI 1 235 Denzity 1.173 Temp. °C. 64.7 796 99.7 43 78 Solubility 3691 40 45 1.285 1 312 Density 1.264

It is insoluble in alcohol Potassium and other ferrocyanides are comoxide, giving the undissociated mercuric cyanide, ferrous hydroxide and potassium bydroxide. They are also decomposed by aurous cyanide or bydroxide, or by finely, divided gold in the presence of oxygen, the complex aurocyanide being formed, as in the cyanide process for dissolving gold (see under Gold Cyanides, p. 486), together with ferric bydroxide (Beutel, Z. anorg. Chem. 1912, 78, 141).

Sodium ferrocyanide,

Na₄Fe(CN)₆,10H₉O,

crystallises in large lemon-yellow monoclinic prisms or rhombohedra, stable in air at the ordinary temperature, but completely debydrated at about 100°C. The dry salt decomposes like the potassium salt on strong heating. The concentrations and densities of saturated solutions have been measured by Farrow (J.C.S. 1926, 129, 49); solubilities are expressed in grams of anhydrous salt per 100 g. of solution:

Temp. °C.	$24.9 \\ 17.11 \\ 1.131$	34·9	49·8
Solubility		20·58	26·20
Density		1·158	1·200
Temp. °C.	64·7	79·6	84·7
Solubility	31·43	36·85	38·15
Density	1·243	1·279	1·292
Temp. °C.	89·6	94·7	99·7
Solubility	38·08	38·25	37·53
Density	1·296	1·287	1·286

A transition point to a lower hydrate occurs at 81.7°C.

Zinc ferrocyanide, Zn₂Fe(CN)₆,6H₂O, is a white amorphous precipitate obtained by the action of calcium or hydrogen ferrocyanide on excess of zinc chloride in aqueous solution. When other soluble ferrocyanides are used, the precipitate contains varying amounts of the other metal (see Treadwell and Chervet, Helv. Chim. Acta, 1923, 6, 559). Reihlen and Zimmermann (Annalen, 1927, 451, 75) postulate the evistence of complex anions containing both zinc and iron, analogous to the copper ferrocyanides and to the Prussian blues. Zinc ferrocyanide is insoluble in dilute acids, completely soluble in alkalis with formation of ferrocyanide and zincate, but only partially decomposed by alkali carbonates.

Hexamethylcarbylamine ferro-salts.—By the action of methyl sulphate on dry potassium ferrocyanide, or of methyl iodide on silver ferrocyanide, E. G. J. Hartley (J.C.S. 1910, 97, 1066, 1725; 1911, 99, 1549; 1912, 101, 705; 1913, 103, 1196) has obtained a series of salts of the general formula [Fe(CNCH₃)₆]R₂, R being a univalent acid radicle and the iron cyanogen complex acting as a kation. On heating with concentrated sulphuric acid, or on boiling with aqueous caustic soda, they are decomposed with quantitativo production of methylamine or methylcarbylamine, showing that all the nitrogen atoms are linked to methyl groups.

Tetramethyl ferrocyanide,

$$\left[\begin{smallmatrix} \operatorname{Fe}(\operatorname{CNCH_3})_4 \\ (\operatorname{CN})_2 \end{smallmatrix}\right]$$

is formed when hexamethylcarbylamine ferrochloride, [Fe(CNCH₃)₆]Ol₂, a representative of the above class of compounds, is heated at 140-150°C. in vacuo for 6-9 hours, when methyl chloride is split off. Hexamethylcarbylamine ferrochloride itself is produced when

$$[Fe(CNCH_3)_6][CH_3SO_4]_2$$

the initial product of the interaction of potassium ferrocyanide with dimethyl sulphate, is boiled with hydrochloric acid or barium chloride. There are two isomeric forms of tetramethyl ferrocyanide, both of which are present in the above product. The a-isomer is much more soluble in water and organie solvents than is the β -isomer, and crystallises with 2 molecules of water of crystallisation, against 6 in the case of the β -isomer. The a-isomer does not dissociate electrolytically in solution, but the β -isomer (according to Hölzl, et al., Monatsb. 1927, 48, 71) is an electrolyte. The a-compound combines with methyl iodide at 100°C, to give hexamethyl-carbylamine ferroiodide, but the β -compound has no reaction. No method of converting one form into the other has been discovered.

Ferricyanides.

TECHNICAL METHODS OF PREFARATION. 1. By Chemical Oxidation of Ferrocyanides.—The commonest oxidising agent is gaseous chlorine, which may react either with solid potassium ferrocyanide, in which case the product is a solid containing potassium chloride, or with a cold 10% solution of ferrocyanide:

$$2K_4Fe(CN)_6+Cl_2=2K_3Fe(CN)_6+2KCl$$

The reaction is stopped when the conversion of the ferrocyanide is just completed, for excess of chlorine destroys the complex radicle. In practice, a small amount of Prussian green is always formed, which spoils the appearance of the product. The difficulty may be avoided by evaporating the solution to the crystallising point and then adding caustic potash to decompose the Prussian green into ferric hydroxide, which can be filtered off, and potassium ferrocyanide, which is apt to contaminate the crystallised ferricyanide. A yield of 85-90% is obtained in the process.

In order to overcome the above objections to the use of chlorine, lead peroxide, calcium plumbate and potassium permanganate have been suggested as alternative oxidising agents, but have achieved no technical importance (Aten, Chem. Weekblad, 1921, 18, 140). More recently, the use of air containing carbon dioxide at a pressure of 80 to 120 atmospheres and a temperature of 80° to 100°C. bas been proposed (Gluud and Keller, U.S.P. 1872929, 1932) for oxidising potassium ferrocyanide solution. W. Klempt (U.S.P. 1908516, 1933) describes a similar process for calcium ferrocyanide solution.

2. By Electrolytic Oxidation of Ferrocyanides.— The electrolysis of ferrocyanide solutions gives rise to ferricyanide and alkali at the anode:

Fe(CN)₆""=Fe(CN)₆""+
$$\bigcirc$$
;
H₂O+ \bigcirc =OH'+ $\frac{1}{2}$ H₂

This method of converting ferrocyanide ruto | The aqueous solution is decomposed by light to ferricyanide has been investigated by Grobe give ferrocyanide. The solution is not as from the electrochemical standpoint (Z. anorg Chem. 1914, 84, 190; Z. Elektrochem. 1914, 20, 334). In practice, the electrolysis of neutral ferrocyanide solutions is carried out at 60°C. in diaphragm cells with nickel electrodes Potassium ferricvanide crystallises out from the anode liquor on cooling, and the mother liquor is replenished with ferrocyanide and returned to the cell. The purpose of using a diaphragm is to prevent reduction of ferra cy anide at the kathode, with consequent decrease of current efficiency. According to Paweck and Hirsch (Z. Elektrochem 1928, 34, 684), the diaphragm can be omitted if an alternating current is superimposed on the direct current

slum ferricyanide, "red prossiate of potash, KaFe(CN), is prepared in the laboratory by the electrolytic oxidation of an alkaline solution of potassium ferrocyanide, or by the technical method of oxidation with chlorine. salt crystallises anhydrous from water in mag nificent, deep red, monoclinio prisms, of density 1.858 (Biltz) The crystals are isomorphous with the cobalti, mangam, and chromicyanides of potassium. The salt is decomposed by 20% aulphuric acid on heating the code of yield hydrocyanic acid, and with 80% acid it gives carbon monoxide and carbon dioxide (Bassett and Corbet, JCS 1924, 125, 1358)

PROPERTIES OF THE FERRICYANIDES -Potas-

The properties of saturated solutions have been measured by Friend and Smirles (JCS. 1028, 2242); the solubilities are expressed in

ams or sait	per 100 g	or solution.	
Temp. °C.	0.1	7.8	15 7
Solubility	23-22	26 06	30 35
Density	_	1 157	1 17
Temp °C.	22 1	28 3	39 9
Solubility	32 08	33 66	37 22
Density	1 187	1,193	1 21:
Temp. °C.	58	81	99
Solubility	41 10	44 70	47 60
Density	1 227		

stable as ferroey anide solution, for on heating with carbon dioxide at 80-100°C. it is appreciably hydrolysed with the formation of pentacyanide. Ferricyanides in caustic potash solution are relatively strong exidising agents, decolourssing indigo instantaneously, and being also reduced by hydrogen peroxide with libera tion of exygen.

Ferric ferricyanide. - When ferricyanide and ferrie salt solutions are mixed a reddish. brown colloidal solution is obtained, which is believed to contain ferric ferricyanide. Haller (Kolloid Z., 1917, 20, 76) examined the solution ultramicroscopically, and showed that the colloidal particles gradually increase in size and a green substance separates out. Reihlen and vou Kummer (Annalen, 1929, 469, 30) produced brown, slightly soluble microcrystals Feg(CN), 2H2O from acid solutions of potassium ferricy anide and ferric mirate. Green ferric ferricyanides are obtained by boiling Prussian blue with concentrated nitric acid, or from potassium ferricyanide and ferric chloride.

Pentacyanides.

As already mentioned (p. 466), the icon heracyanides can undergo a decomposition of the complex by which one ionic cyanide group is replaced by a molecular group, giving a new pentacyanide complex with a valency lower than that of the original complex The most important transformations of hexacyanides into

pentacyanides are listed in the table below. Other substituents which may be found in complex pentacyanides are NO, AsO, SO,

NH, and substituted ammonias. The correct constitution of the nitroprussides was first given by Gerhardt (Traité de chimis organique, 1853, I, 344). K. A. Hofmann carried out extensive work on their reactions (Z. anorg. Chem 1893, 10, 262; 1896, 11, 31, 278; 1896, 12, 146), and was able to

explain his observations on the hasis of the co-

	Ferroc) anide [Fe(CN)4]"" gives	Perricyanide [Fe(CN),]" gives
By warming aqueous solutions, action of light or acids By action of strong exclaining agents are additions of additions of additions with carbon monorate solutions with carbon by warming with mtric or nutrous acid	squopentacyanoferroate, [Fe(CN) ₈ H ₂ O]"" carbonylferrocyanide, [Fe(CN) ₂ CO]"" nitrosoprussuate, IFe(CN) ₈ NO]" (mtroprosside)	aquopentacyanofernate, [Fe(CN) ₅ H ₂ O]" carbonylfernocyanide, [Fe(CN) ₅ CO]" nitrosoprussate, [Fe(CN) ₅ NO]" (nitroprusside)

ordination theory (Hofmann, Annalen, 1900, is formed by the action of carbon monoxide 312, 1). Other important work on the transiton of hexacyandes into pentacyandes is described by Manchot (Ber. 1912, 45, 2869), Cambi (Cazetta, 1911, [1, 41, 157) and Jimori (Z. anorg. Chem. 1927, 167, 145) Polassium carbonylferrocyanide,

K,[Fe(CN),CO]H,O,

on an aqueous solution of potassium ferro cyanide at 135°C. in a scaled tube (Muller, Ann chim. phys. 1889, [vi], 17, 94): K_Fe(CN),+CO+2H,O

=K₃[Fe(CN),CO]+NH₃+HCOOK

It may also be made by heating potassium ferrocyanide with sulphuric acid until the evolu-

Carbonylferrocyanides are present in the cyanide mud and spent oxide of the purification plant of gas works; according to Colman (Analyst, 1908, 33, 261), from 2 to 5% of the cyanogen (and sometimes as much as 20%) is present in this form. When cyanide mud and spent oxide are extracted with caustic soda, and the extract mixed with 5 times its volume of methylated spirit, the sodium ferrocyanide crystallises out completely in a few hours, after which the carbonylferrocyanide can be pre-cipitated as the amorphous, deep violet ferric salt by adding ferric chloride to the acidified solution. The deliquescent calcium salt, which crystallises with 8 molecules of water, is obtained by boiling the ferric salt with milk of lime. The copper salt, a green precipitate soluble in excess of ammonia, is characteristic. The carbonylferrocyanides resemble the corresponding ferrocyanides, but are paler in colour, more soluble in water, and more stable towards oxidising agents.

Sodium nitroprusside,

Na₂[Fe(CN)₅NO]2H₂O,

is made by the action of dilute nitric acid on a ferrocyanide solution (Playfair, Phil. Mag. 1850 [3], 36, 197). The solution is neutralised with sodium carbonate and evaporated to crystallisc out potassium nitrate, which is separated before finally crystallising the nitroprusside. The mechanism of its formation is that the nitric acid oxidises the ferrocyanide to ferricyanide, itself being reduced to nitric oxide, and the two products then react in accordance with the equation:

$$Fe(CN)_6"'+NO+H^+$$

$$=[Fe(CN)_5NO]"+HCN$$

The salt can also be formed from solutions of ferrous sulphate, sodium nitrite and potassium cyanide. It crystallises in dark red prisms, soluble in 2½ parts of water at 16°C. The solution is not precipitated by a ferric salt, and gives a deep violet coloration with alkali sulphides.

Sodium aquopentacyanoferroate,

$$Na_3[Fe(CN)_5H_2O]5H_2O$$
,

is obtained by the action of reducing agents such as phenylhydrazine or hydroxylamine, or of oxidising agents such as potassium hypobromite or hydrogen peroxide, on sodium nitroprusside solution below 0°C. The NO group of the nitroprusside is replaced by water, and the iron of the complex is converted from FeIII to FeII. In the latter respect, the action of hydrogen peroxide is analogous to its behaviour with ferricyanide, which it reduces to ferrocyanide. The salt crystallises in reddish-yellow needles, giving an intensely yellow solution which does not yield a violet coloration with alkali sulphides.

Sodium aquopentacyanoferriate,

 $Na_2[Fe(CN)_5H_2O]$,

tion of carbon monoxide begins, and then is obtained by oxidising the ferroate com-cooling (see Hölzl, Monatsh. 1930, 56, 79). pound (described above) with bromine water. Carbonylferrocyanides are present in the It is a deep violet-blue substance. The corresponding potassium salt is obtained by the action of excess chlorine on potassium ferricyanide, and is identical, according to Cambi (Gazzetta, 1911, 41, I, 57), with the "potassium perferricyanide" obtained by Bong (Bull. Soc. chim. 1875, [ii], 24, 268) and Skraup (Annalen, 1877, 189, 368) by the action of potassium chlorate on potassium formations of the contraction of t potassium chlorate on potassium ferricyanide solution.

CONSTITUTION OF THE COMPLEX IRON CYAN-IDES.—According to Werner's co-ordination theory ("Neuerc Anschauungen auf dem Gebiete der anorganischen Chemie," 5th ed., 1923), complex salts are to be regarded as built up by the "co-ordination," through co-valent, non-ionisable linkages, of a number of atomic groups round a central atom, giving a more or less stable nucleus. According to the nature of its component parts this nucleus may or may not be able to combine with other atoms by means of electrovalent, ionisable links. The significance of the square brackets in the usual method of writing Wernerian formulæ is that all the linkages between groups within the brackets are non-ionisable. The co-ordination number of the central atom bears no obvious relation to its valency; the maximum co-ordination number of carbon, nitrogen and boron is 4, of the great majority of elements is 6, and of some of the heavier elements (e.g. molybdenum and tungsten) is 8.

The great stability of the ferrocyanides and

ferricyanides has already been mentioned. The first decomposition of the complex ions is the removal of one cyanogen ion, which is always replaced by another molecular group, such as H_2O , NO, or CO, giving a new complex in which the iron atom is still associated with six groups. Thus the whole series of complex iron cyanides, derived from divalent and trivalent iron, provides an example of co-ordinated compounds in which the iron (in either valency state) has a well-defined co-ordination number of 6. The following list of compounds illustrates the way in which the electrovalency of the 6co-ordinated complex is altered by the abstraction of one or more cyanide ions, to be replaced by the same number of neutral atomic groups:

* The corresponding ferricyanide derivative is wanting (Hartley, J.C.S. 1914, 105, 521).

In the case of a complex containing six co-ordinated groups, Werner held that the six groups are arranged symmetrically at the corners of a regular octahedron, the co-ordinating atom being at the centre. Hence complexes of the general formulæ Ma_4b_2 and Ma_3b_3 should 478

for the second. Hartley's two moment tetramethyl ferrocyanides (p. 475) are regarded as stereosomers by Glasstone (J.C.S. 1930, 321), and the same may be true of his isomeric trimethyl cobalticyanides (ibid. 1914, 105, 521).

On the other hand, no isomerism would be expected in the simple ferro and ferri-cyanides. Briggs (soid 1911, 99, 1019) described α - and β isomers of alkali ferrocyanides with slight differences in physical properties, but Bennett (sold 1917, 111, 490) showed that the two forms are crystallographically identical, and Piutti (Ber. 1912, 45, 1830) and Getman (J. Physical Chem 1921, 25, 147) found no differences in the absorption spectra Locke and Edwards (J. Amer. Chem. Soc. 1899, 21, 193, 413) described the conversion of ordinary red e potassium ferricyanide into an isomeric green β modification by treatment with acids. The green product had a different crystalline form, a hsorption apectrum (Getman, I c), and chemical properties (Bellucei and Sahatmi, Atta R Accad Linces, 1911, [V], 20, I, 176). The view is now held by Briggs (JCS, 1920, 117, 1026; J. Physical Chem 1928, 32, 1422) that no isomerism exists in the alkali ferro. and ferri cyanides, the observed differences in the properties of the \$\beta\$ forms being die, in the case of ferrocyanide, to the prosence of small amounts of aquopentacyanoferroate, K, [Fe(CN), H,O], and in the case of formcyanide, to the existence of a double salt between ferricyanide and a quo pentacyanoferriate having the formula

It is a shortcoming of Werner's theory that his structural formulæ lack the definiteness of those of organio chemistry, and that the existence is implied of a new kind of valency for the coordinating atom, having no relation to its normal electrovalency or covalency. The difficulty has been dispelled by the application of the electronic mechanism of valency to Fleck (" Die Fabrikation chemischer Produkte tion of the electronic mechanism of vaccary to several the resolution measurement around or ordination (folgorick, J. O. 8 1923, 123, 725; laws terractione Abfallen, 1862) and Chaster "The Electronic Theory of Valency," 1921). (B.F. 15911 of 1891) proposed the use of charcoal According to the electronic interpretation, a to reduce the cyanate, but it appears unlikely covalent, non-ionisable linkage is formed by the that this reduction would occur at the relatively sharing of two electrons between the combining low temperatures used in the fusion, atoma, each electron simultaneously forming a part of the stable electron configurations of both atoms. In the normal covalent link, one shared electron is provided by each of the atoms, but in the co ordinated link both are supplied by one of the atoms, termed the "donor" Henco a co ordination number of 4 corresponds to the formation of a stable octet of electrons, and of 6 to the formation of a ring of twelve, another stable configuration

exist in the two stereoisomeric forms repre-aented by:

gives a satisfactory representation of the pro-perties of the simple ferro- and ferri-cyanides of the alkalı and alkalıne earth metals, difficulties arise in the case of the heavy metal salts such as Prussan blue, copper ferrocyanide, and rine ferrocyanide, which have been met by the assumption of a complex having two nuclear atoms, eg. Fell and Felli in the case of Prussian blue (see under Constitution of the Blue Compounds, p. 473), Zn and Fe¹¹ in the case of zine ferrocyanide. In these cases the co ordination number of each of the nuclear metallic atoms sa probably 4. Similar bi nuclear complexes may exist in the numerous double salts formed by ferrocyanides, the less electropositive of the two metals tending to enter the ferroeyanido complex. In this connection it is of interest to note that Berkeley, Hartley and Stephenson (Phil. Trans. 1909, 209,A, 319) conclude, from measurements of the osmotic pressure and electrical conductivity of solutions of ferrocyanides of calcium and stroptium, that these salts give in solution undissociated mole cules of twice the simple formula weights, indicating a tendency to similar binuclear complex formation even in the alkaline earth ferrocyamdes.

H. ALKALI METAL CYANIDES.

TECHNICAL METHODS OF PREPARATION -1. From Ferrocyanides The earliest method of preparing potassium cyanide from ferrocyanide, due to Berzelius, consisted in heating dry potassium ferrocyanide to redness;

K_Fe(CN),=4KCN+Fe+2C+N.

A 33% loss of cyanogen occurs in this reaction, which can be reduced by adding potassium carhonate so that the fusion is estried out with a muture of 8 parts of ferrocyanide to 3 parts of potash (F. and E. Rodgers, Phil, Vag. 1834, [3], 4, 91). The melt is allowed to and until the iron and carbon have separated out, and is then poured into moulds, giving white, porcelain like cakes containing from 30 to 70% of KCN, This product was used until about 1890 for electroplating and other purposes. Liebig showed that eyanato was formed during the reaction, thus:

 $K_4Fe(CN)_4+K_2CO_2$ =5KCN+KCNO+Fe+CO₂

Erlenmeyer (Ber. 1876, 9, 1840) described a much more advantageous process, in whi h a pure product containing approximately 40% CN (equivalent to 100% KCN) was obtained by fusing dry potassium ferrocyanide with metallic sodium in the proportions required by tbe equation .

K4Fe(CN)4+2Na=4KCN+2NaCN+Fe

The process became technically important in the While the concept of 6-co ordinated iron hands of the Deuts. Gold u. Silber Scheide-

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anstalt vorm. Roessler, V. Intern. Kongress für Angewandte Chemie, Berlin, 1903, I, 638) when sodium hecame a commercial product in 1890. The spongy iron is filtered off from the melt and then pressed out while still red-hot; filtered cyanide is cast in cakes and gives a very pure commercial product. Since 1905, sodium ferrocyanide has been used instead of the potassium salt, and the product is almost pure sodium cyanide. In England the surplus of ferrocyanide from gas works over that required in the colour industries was formerly converted into cyanide by this process. Modifications have heen described by Hetherington, Muspratt and Hurter (B.P. 5832 of 1894), who used a sodiumlead alloy instead of sodium and obtained a three-layer product, the powdered iron floating hetween the upper liquid cyanide and the lower liquid lead, and by Crowther and Rossiter (B.P. 9275 of 1894), who used zinc alone for reducing the ferrocyanide, subsequently decomposing the zinc cyanide hy fusion with alkali carbonate. Neither modification has achieved practical importance.

From Ammonia and Alkali Carbonates.— Two important processes which have been large producers of cyanide, utilising the same reactions but under considerably different working conditions, fall under this head. In Siepermann's process (G.P. 38012 of 1886; B.P. 13697 of 1889, 9350 and 9351 of 1900) a dry mixture of 2 parts of wood charcoal and 1 part of potassium carbonate is introduced into the upper end of a vertical iron retort, and is treated at a dull red heat with dry ammonia gas introduced at a point about half-way down the length of the retort. Absorption of the ammonia occurs with production of potassium cyanate, probably through the intermediate formation of potassium carbamate (according to Pfleger, see Deuts. Gold-u.-Silber-Scheideanstalt vorm. Roessler, V. Intern. Kongress für Angewandte Chemie, Berlin, 1903, I, 638):

$CO(OK)_2+NH_3=CO(NH_2)OK+KOH$ CO(NH₉)OK=KCNO+H₉O

The product then falls into the lower part of the retort, which is maintained at a hright red heat. At the higher temperature the cyanate is converted into cyanide, and carbon monoxide is produced which escapes through the upper part of the retort with the hydrogen formed hy the decomposition of some of the ammonia. According to Pfleger (l.c.) the conversion of the cyanate proceeds in two stages:

2KCNO=K₂CN₂+CO₂

(cf. Drechsel, J. pr. Chem. 1877, [ii], 16, 201, on a similar decomposition of alkaline earth cyanates), and

 $CO_2+C=2CO$.

Some of the potassium cyanamide may decom-

The product of Siepermann's process is allowed to cool out of contact with air, and is then systematically leached with exclusion of air (B.P. 9351 of 1900). Potassium cyanide may he precipitated from the extract by adding excess of potassium carhonate, or better hy evaporating nearly to dryness and extracting unchanged potassium carhonate with a minimum quantity potassium extronate with a infiliation question of water at 66°C. The residue, containing potassium cyanide and some unconverted cyanate, is treated with refrigerated water (between -18° and +5°C.) to dissolve the cyanide and leave the cyanate, and the cyanide is crystallised as a damp, deliquescent mass (B.P. 9352 of 1900). This product had to he fused with cyanide produced from ferrocyanide in order to make it suitable for export to the gold

The second process of this kind is the Beilby process (B.P. 4820 of 1891), which first became a producer in 1892, the same year as Siepermann's. Gaseous ammonia is forced through molten potassium carhonate, and the addition of charcoal is made gradually, the total quantity added heing hardly more than that theoretically required by the equation:

$$K_2CO_3+4C+2NH_3=2KCN+3CO+3H_2$$

In this way the leaching step of Siepermann's method is avoided, for it is possible to run off the molten mass from the retorts, through filters for the small amount of unreacted charcoal, directly into moulds to give a white, saleable product. In practice, previously mado cyamde was added to the potassium carbonate on starting the process, in order to lower the melting-point of the reaction mass, and the passage of ammonia was continued until an analysis of the melt showed that the reaction was The reaction temperature was 900-1000°C., and the formation of cyanate and its conversion into cyanide occur simultaneously.

As regards the relative merits of the two pro-Siepermann's gives higher yields of eyanide based on the ammonia used because, at the lower temperature of the first stage reaction, less decomposition of ammonia into its elements occurs. The higher temperature in Beilby's process is necessary, however, for the maintenance of a reasonably high rate of reaction since hy working in a medium of molten salt the advantage of the large surface exposed hy Siepermann's dry mass of alkalised charcoal is sacrificed. The dissociation of the ammonia in Beilhy's process is minimised by introducing the gas into the retort through tubes lined with copper, this metal having no catalytic effect on the decomposition of ammonia. The yields on ammonia actually obtained are 80-85% in Siepermann's, and 55-60% in Beilhy's process. According to Readman (J.S.C.I. 1889, 8, 757) it is possible to obtain an almost quantitative Drechsel, J. pr. Chem. 1880, [ii], 21, 77) ; finally: yield hy working at a low enough temperature. The great advantage of Beilby's process is the easy working up of the product into a readily marketable form.

3. From Ammonia and Alkali Metals.—The pose directly into cyanide, potassium and nitrogen, as this reaction is known to occur below 800°C. in the absence of excess charcoal. In 1894 (B.P. 12219 of 1894; G.P. 90999, F.P. 239643), and rapidly became the most important plant was started at Glasgow in 1900, and a process in use for cyanide mannfacture. The complete reaction is represented by the equa-

According to Castner's earlier claims (BP. 21732 of 1894), the reaction is best carried out in two stages, in the first of which liquid sodium is allowed to react with dry ammonia gas at 300-400°C, in a horizontal cast iron retort fitted with a series of vertical baffles to increase the length of passage of the gas over the liquid Sodamide is formed, which is stable up to temperatures approaching 500°C. (Titherley, JCS. 1894, 65, 504):

In the second stage the molten sodamide is run on to red-het wood charceal in a vertical cylindrical retort Carbon absorption occurs with the formation of sodium cyanamide, a compound which is relatively etable up to 600°C

Cyanamide may also be formed, according to Pfleger, by the interaction of sodamide and sodium eyanide already formed

The second stage of the Castner process 12 completed by raising the temperature of the eyanis ing furnace to 700-800°C, when the eyanamide melt absorbs further quantities of carbon to form molten evanide:

According to G.P. 148046 (Deutz, Gold u-Silber-Scheideanstalt vorm. Roessler), it is possible to carry out the whole process in one retort, the sodamide being produced at the lowest possible temperature, and the temperature then gradually raised for the succeeding atages of the reaction. The molten charge is run off from the retort through filters to remove small amounts of solid impurities (such as unreacted charcoal), and is east in iron moulds to give purs white cakes with a crystalline fracture, and analysing 97-98% NaCN, 21% Na₂CO₂, 03% NaCNO, and 002% Na₂S Instead of charcoal, velatile organic com-

pounds such as acetylene (G.P. 149678) or tramethylamine (G P. 223027) may be added to the melt as sources of carbon, but such variations have not come into general practice The source of the ammenia used in the operation of the Castner process was formerly the by-product of coal distillation, but since 1918 there has been a considerable change over to the use of aynthetic ammonia. This is reflected in H. Freudenberg's B P. 265639, 1926, which claims the collection and recombination with nitrogen of the hydrogen produced in the Castner cyangle synthesis. This hydrogen is relatively pure. containing only amall quantities of methane and

The Castner process was first put into opera-

third near New York in 1901. Since that time it has continued to provide the greater part of the world's cyanide requirements, and still retains its pre-emment position.

4. From Thiocyanates .- The usual method of preparing cyanides from thiocyanates is first to convert the thiocyanate into ferrocyanide by one of the methods already described (p. 469), and then to work up the ferrocyanide into cyanide by the Erlenmeyer method as described on p. 478. On the other hand, the direct desulphurisation of thiocyanates can be carried out by fusion with lead or zinc (Warren, Chem. News, 1890, 62, 252; Playfair, B P. 7764 of 1890) The fusion is carried out at 400°C, and gives a 70-80% conversion to cyanide (Playfair, J S C I. 1892, 11, 14; Conroy, ibid, 1896, 15, 8), which is extracted from the ecoled melt with water, Playfair (le) also found that hydrogen reduces potassium thiocyanate at a dull red heat, giving potassium cyanide, potassium sulphide, and aulphuretted hydrogen, Conroy, Heslop, and Shores (J S C.I. 1901, 20, 320) found that, on reduction with hydrogen, about 70% of the nitrogen is obtained as potassium cyanide and 20% is further reduced to hydrocyanic acid An interesting proposal for the direct desulphure sation of ammonium thiocyanate by heating it with finely divided copper at 300-500°C. has recently been made by Konig and Siegle and Co (B P 438358 and 438402, 1934). Cuprous sulphide is formed and the vapours of ammonium eyanide are evolved and are absorbed in water to give a dilute solution. It is not possible to separate solid ammonium eyanide from this solution, which is highly unstable on keeping, hut it is claimed that the solution is suitable for use in the extraction of gold from its ores

Zine oxide may be used as a desulphurising agent if carbon monoxids is also present According to G P. 588823, 1032 (N.V. Stixstof bindingsindustrie, Nederland), a 89% yield of sodium eyanido is obtained by beating sodium thiocyanate above 400°C. with zine oxide and a gas containing carbon monoxide :

Lime or chalk may he used instead of zine oxide (B P. 401627, 1932); in this case the initial crude product contains a high percentage of sodium cyanate, which can be reduced to cyanide, however, by continuing the passage of carbon monexide:

NaCNS+CaO=NaCNO+CaS NaCNO+CO-NaCN+CO,

5 From Hydrocyanic Acid -The recovery of the hydrocyanic acid content of "Schlempe" gases (see Hydrocyanic Acid, p. 493), in the form of solid sodium cyanido is an important technical process. The cyanised gas, which contains 7% HCN, 7% NH₃, and 24% CO₄ hy volume, as freed from ammonia by acrubbing with diinte oulphuric acid, and separated from carbon dio vide and enriched in hydrocyanic acid by scrubbing with water and fractionally distilling the aqueous HCN-extract. The gas contion at Frankfurt-on-Main in 1899, a second taning the concentrated HCN thus obtained 15

strength that a saturated solution of sodium cyanide (45% by weight) is produced (Muhlert, Chem. App. 1925, 12, 156). This solution is evaporated in vacuo to a paste, which is centrifuged to give a product containing 10% of water, from which the rest of the water is removed by drying on heated trays in vacuo. The dried powder is then briquetted (B.P. 18096 of 1907), and the product appears in commerce with the following analysis: NaCN 92-93%, Na₂CO₃ 3.5%, NaCNO 0.7%, NaCl 1.0%, NaOH 0.5%, Na₂SO₄ 0.2%, Na₂S 0.02%.

In the above process, prior removal of carhon dioxide from the gas is necessary in order to avoid excessive formation of carbonate during the neutralisation. In carrying out this step, care must be taken that the hydrocyanic acid is not present in excess, because of the tendency of free hydrocyanic acid to polymerise in contact with alkaline substances such as cyanides. If this polymerisation occurs, the product is discoloured and its commercial value impaired. The evaporation of the cyanide solution must be carried out in vacuo at a relatively low temperature in order to minimise saponification of the cyanide to formate and ammonia, which results in loss of cyanide efficiency and in formate contamination of the expedients have product. Various suggested to avoid the evaporation step, hut none has found practical application. Interest in the problem has, however, heen revived by the development of the newer cyanide syntheses by high-temperature gas reactions and hy formamide dehydration, which reactions yield their hydrocyanic acid in the form of a dilute gaseous mixture similar to the cyanised "Schlempe"

According to G.P. 160637, 1903, the gas containing the HCN free from carbon dioxide may be passed over solid flake caustic soda at 200°C., so that the water produced (by the reaction NaOH+HCN=NaCN+H₂O) is vaporised, hut no melting of the solid occurs. It is necessary in this method to grind the product when it contains 50-60% NaCN, in order to expose fresh surfaces of caustic soda to the gas. Wiggers (Z. anal. Chem. 1876, 15, 448) obtained pure potassium cyanide by reacting hydrocyanic acid with a solution of caustic potash (3 parts) in 95% alcohol (1 part), washing the potassium cyanide precipitate with alcohol and drying it over sulphuric acid. Guignard and Watrigant (G.P. 243469, 1911) used alcoholic caustic soda for the preparation of sodium cyanide, and added metallic sodium from time to time to react with the water produced, so as to prevent the saponification of hydrocyanic acid to ammonia and formic acid.

The evaporation of aqueous cyanide solutions may also be avoided hy salting out the cyanide from solution with solid alkali (Tscherniac, G.P. 182774, 1906; Chemische Fabrik Schlempe, G.P. 192884, 1906), the salting liquor being used again in the absorption. The Stassfurt Chemicals III is the salting liquor being used ische Fabrik (G.P. 129863, 1901) claims that an anhydrous cyanide containing 43.5% CN can be The first attempts to carry out the reaction obtained by crystallising from a solution on a technical scale were made by Possoz and

absorbed in aqueous caustic soda of such a containing the sodium and potassium salts in 2KCN:NaCN, whereas sodium the ratio cyanide gives the dihydrate.

A considerable amount of work has been done on the interaction of gaseous hydrocyanic acid with solid sodium carhonate to give sodium cyanide:

$Na_{s}CO_{3}+2HCN \rightleftharpoons 2NaCN+H_{2}O+CO_{2}$

In the earlier experiments, high temperatures (700-900°C.) were used, and the cyanide was the run from the reaction chamber in molten state (Roeder and Grünwald, G.P. 134102, 1901. Tscherniac (G.P. 145748, 1902) worked at 450°C., and claimed to produce sodium cyanide of 98-99% quality. More recently attention has been paid to the physical form of the carbonate used, which greatly affects the speed of reaction and the strength of cyanide attainable in the solid product. E. Hene (B.P. 375636, 1931) recommends the use of the light form of sodium carhonate produced hy calcining hicarbonate 500°C., and conducts the absorption of hydrocyanic acid at 360°C. The Gesellschaft für Kohlentechnik (F.P. 751191, 1932) describes the production of similarly reactive carbonate hy dehydration of the crystalline hydrates under temperature conditions such that no solution, fusion, or fritting of the salt occurs. The HCNfixation reaction is reversible, and the equilibrium

$k=[HCN]^2/[H_2O][CO_2]$

precludes high percentage conversions of gaseous hydrocyanic acid, unless one or both of the gaseous reaction products, steam and carhon dioxide, are removed from the system. Thus A. D. Macallum (U.S.P. 1966253, 1934) describes the conversion of the hydrocyanic acid in the gaseous product of formamide dehydration (see Hydrocyanio acid, p. 495). into solid sodium cyanide by passing the gas over reactive sodium carbonate at 200-500°C., removing either or hoth of the gaseous hy-products, water and carbon dioxide, and recirculating the residual gas.

6. From Nitrogen and Alkali Metal Carbonates. -Of the syntheses of alkali metal cyanides from elementary nitrogen the one which has achieved the greatest practical importance is the action of nitrogen on a mixture of alkali carbonate and carbon, generally in the presence of a metallic catalyst. The reaction is expressed by the equation:

$$Na_2CO_3+4C+N_2=2NaCN+3CO$$

Such a synthesis may have heen carried out hy Scheele in 1783 (see "Chemical Essays," 1901, reprint, p. 288) when he noticed the formation of ferrocyanide on fusing charcoal with alkalis, and was certainly used by L. Thompson (Mechanics Magazine, 1839, 31, 92) when he substituted atmospheric nitrogen for nitrogenous animal matter in the preparation of ferrocyanides. Thompson realised the importance of the presence of iron, which lowers the temperature necessary for eyanide formation.

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Boissière (B P. 9985 of 1843) in France, and an | added. The open grained atructure required for account of further large scale experiments in easy access of nitrogen is obtained by briquetting England is given by Graham (Report of Juries, Exhib, of 1851, 1, 95) Wood charcoal was soaked in a solution of patassium carbonato and dried to give a mixture containing about 20% K.O, which was heated to a white heat (probably 1,300-1,400°C) in fire-brick vertical retorts. In order to supply as rapidly as possible the large quantities of heat necessary to attain this high reaction temperature, and te make good the heat absorbed by the ende thermic reaction (amounting to 645 cals, per g mol of potassium cyanido produced), hot furnaco gases (consisting of nitrogen and carbon dioxide) were drawn directly the cyanising mass through alots in the brick work of the retort by means of a suction applied at its upper end Abovo the high temperature, eyanising acction of the retort was a section in which the preliminary drying of the alkalised charcoal by the hat furnace gases took place, and below it was a cooling section from which the cyanised product was withdrawn into ferrous salt solution for the production of ferro-cyanide The process failed financially owing to high potash losses and the short life of the retorts About one third of the alkali and onehalf of the charcoal were eyanised

Victor Alder of Vienna clearly recognised the three important conditions which contribute to the successful operation of this process, namely, the catalytic action of iran (B P. 1001 of 1880), the intimata admixture of the solid reactants, and the porosity of the charge as as to admit of free access of nitrogen In B P. 5617 of 1881 he describes the preparation of the charge by mixing charcoal with alkali and adding sawdust or other material, so that on heating the mixture a porous mass results. The catalytic metal may be added as such, or as oxide, ar as a salt solution which will hydrolyse on heating to give a deposit a current of nitrogen, and gave a praduct con of the oxide, which in turn is reduced to the taining 152% NaCN, carrespanding to a metal under the cyanising conditions catalytic activity of iron (and to a less extent of other metals) was studied by Tauber (Ber 1899, 32, 3150; Chem. Ind. 1903, 26, 26) average conversion of alkali to cyanide in these experiments was anly 10%, and the maximum

Great progress in this cyanide synthesis was shown by the work of J. E. Bucher in America porous, highly reactive alkalised coke for the (Ind Eng. Chem 1017, 9, 233, BP 277) reaction. It is claimed that very porous and of 1912, 13332 and 23,292 of 1914, 2286 of active briguattes are made by mixing sedium 1915), with the result that the process was bicarbonata with coke and a small amount of commercially developed in that country drams the later years of the War of 1914-18 Descriptions of small, acms technical and full scale plant 1939; Bartling, U.S.P. 1937129, 1931; the appear in the literature, and in aildition to the appear in the interactive, and in similation to the evolution of except adjusted from the olderboards article by Bucher mentioned above, the following the proposal in the mass the necessary promety. A matter proposal in the start proposal in (which, however, gives inferior results) and presence of the alkali confers upon the coke an iron oxide (Fc,O3 or Fe,O4) are separately enhanced reactivity. It is claimed for activated ground to pass a 100 mesh sieve, then re- masses of these types that they can be cyanised ground together and dry sodium carbonate with mitrogen at temperatures as low as 830-

the mixture : the powder is kneaded at 100°C with water sufficient to form the monehydrate Na CO, H.O, and the paste extruded in the form of rods 1 in, long and 1 in, in diameter, which are dried immediately to prevent decahydrate formation. The proportions of carbon, iron, and sedium carbonate in the mixture aro from 2:2:1 to 2:2:2, the greatest reaction velocity being obtained with the lowest sodium carbonate content. The reaction temperature is 950-1,000°C, at which the briquettes are slighty plastic: reaction does not occur below 700°C., and temperatures above 1,150°C. (the cutectic of iron-carbon) should be avoided, because fusion of the iron particles diminishes there active surface. The method of heating may be externally by gas or electric power, or internally by using the bot charge itself as an electrical resistance. This last method is preferred by Bucher, surro it affords a means of supplying the necessary heat (including that of the endothermic reaction) at a high rate. Pure nitrogen or producer gas may be used as the source of nitrogen, According to De Kay Thompson (i.e.), the results with producer gas are inferior, and this may be due to the reversibility of the reaction .

Na,CO,+4C+N, =2NaCN+3CO-138 5 kg -csls

The use of producer gas decreases the partial pressure of nitrogen and increases that of carbon monoxide, displacing the equilibrium to the left, De Kay Thompson states that the amount of nitrogen required for satisfactary results is three times the theoretical

In a typical batch experiment, a carbon-ironcarbonate mixture of the composition 2:2:1 was heated in 13 minutes fram 710° to 920°C, m The | 92% conversion of carbonata into cyanida. At 900-930°C, aome volatilisation af sodium examide occurs, so that the upper layers of a batch product are richer in cyanide than the

lower ones. The more recent work on this process for eyanido synthesis has been directed towards the development of methods of preparing masses of evolution of carbon dioxide from the bicarbonste

that the Bucher process gives better results when the carbon and iron in the reaction mixture are in excess; A. Mentzel finds for his noncatalysed cyanising process that it is advantageous to use a 25-33% excess of carbon, the excess being added in the form of granules of diameter not more than 3 mm., to provide porosity and contact surface (G.P. 614100, 1930).

The fact that the cyanising reaction is reversible was proved by the experiments of Ferguson and Manning (Ind. Eng. Chem. 1919, 11, 946). These authors heated mixtures of charcoal (1 part), iron (1 part), and sodium cyanide or carbonate (3 parts) with a range of definite mixtures of nitrogen and carbon monoxide. The composition of the product was independent of the nature of the solid raw materials, but varied with the temperature and the composition of the gas. At 946°C a gas containing 5% carbon monoxide gave a solid product with 75% of the alkali cyanised, and one containing 48% CO a product with only 27.5% cyanised. At 1,000°C. the first gas mixture gave 80% cyanisation, and the second 59%. Ingold and Wilson (J.C.S. 1922, 121, 2278) took precautions to avoid errors due to the volatilisation of sodium cyanide at these high temperatures, and found that, in the range 850-950°C., the relation between the composition of the liquid phase (Na₂CO₃ and NaCN) and the gaseous phase (CO and N₂) in equilibrium can be expressed by

$$\log\left(\frac{x^2}{1-x} \cdot \frac{(1-p)^2}{p} \cdot \pi^2\right) = 23.91 - \frac{31,180}{T},$$

where x is the fraction of the sodium which is present as cyanide, p is the mol. fraction of nitrogen in the gas, and π is the total pressure.

The Bucher process was operated on a technical scale for a short time during 1918 by the Government of the United States, whose plant is described by Brown (Ind. Eng. Chem. 1919, 11, 1010). The retorts were stationary nichromo tubes heated by producer gas, each tube having a life of 11-14 days and producing about 7½ lb./hour of sodium cyanide. Pure nitrogen from an air-liquefaction plant was used for the cyanising. The recovery of cyanide from the product by lixiviation with water is attended with the danger of ferrocyanide formation, though Chickering (Chem. Met. Eng. 1924, 31, 967) describes such a method of obtaining anhydrous sodium cyanide from the via the dihydrate. The method adopted at the plant in question was leaching with liquid ammonia under pressure. By this method unchanged alkali and carbon are left behind and can be reground with fresh raw materials for return to the retorts. Sodium cyanide of 92% purity was obtained from the liquid ammonia solution in the form of a white powder by evaporating off the ammonia in stcam-heated evaporators, the ammonia loss being relatively low at 2 lb. per 100 lb. of cyanide made. A related process was worked between 1923 and 1927 by the California Cyanide Co. of Los Angeles. The retorts, made of nichrome or other nickel chromium iron alloy (B.P. 199900,

850°C., apparently in the absence of iron | 15 in. in diameter and having a length of catalyst. It has already been mentioned | 10 ft. heated to 950-1,050°C. (B.P. 202383, 1922). The charge consisted of a finely ground and granulated mixture of 54 parts of coke, 44 of sodium carbonate and 2 of iron (B.P. 136750, 1919), which passed down the tube in countercurrent to the flow of nitrogen and was converted into a product containing 25-30% NaCN. As the primary object of this manufacture was the production of hydrocyanic acid. the solid product was treated with moist carbon dioxide (U.S.P. 1385335, 1385336, 1921), which was passed through the solid spread out on perforated trays:

2NaCN+H₂O+CO₂=Na₂CO₃+2HCN

In carrying out this reaction it is necessary to diminish the time of contact of the hydrocyanic acid with alkali, in order to minimise its decomposition and polymerisation. The product gases were dried and the hydrocyanic acid condensed by cooling. If desired, the dry hydrocyanic acid may be converted into sodium cyanide by reaction with sodium carbonate as described under Alkali Metal Cyanides,

p. 481. 7. From Nitrogen, Alkali Metals, and Carbon. The direct synthesis of potassium cyanide from its elements was first observed by Marchand (J. pr. Chem. 1850, 49, 351) in testing for the presence of nitrogen in cast iron and steel by Lassaigne's method of heating the sample to red heat with potassium, extracting the product with water, and applying the Prussian blue test to the solution. Marchand found that, by repeated treatment of a sample of powdered iron, he was unable to exhaust the cyanide-forming reaction, and he showed that the cyanide was derived from gaseous nitrogen by carrying out the reaction in a closed tube containing nitrogen, which was completely absorbed. Since he was unable to prepare Prussian blue from the mixture of iron and carbon obtained by heating ferric succinate or benzoate, Marchand concluded that the carbon must be in chemical combination with the iron. At a later date, however, Remsen (Amer. Chem. J. 1881, 3, 134) showed that iron powder loses its catalytic activity when superficially oxidised by exposure to air, and this may explain Marchand's failure with his simple mixtures of carbon and iron. observations were made by Tauber (Ber. 1899, 32, 3150).

Bucher in 1912 made use of the catalytic activity of iron in the synthesis of sodium cyanide from its elements (B.P. 11797 of 1913: U.S.P. 1082845, 1913; 1116559, 1914). The iron and the carbon are intimately mixed, and reaction with sodium and nitrogen proceeds at 500-1100°C. Bucher, like Marchand, believes that the iron acts as a solvent for the carbon, which is then capable of entering into reaction with sodium and nitrogen. In the cyanide synthesis from sodium carbonate (see previous section), Bucher postulates the initial formation from the carbonate and carbon of some metallic sodium, which is continually consumed by conversion into cyanide. This hypothesis makes 1922), were slightly inclined rotating tubes, the rate of cyanide formation dependent on the rate of solution of solid carbon in solid iron and, 161, [B], 670). Potassium cyanide reacts with in view of the extremely rapid formation of tetrathionate in aqueous solution to form in view of the extremely rapid formation of cyanide actually observed, is bardly acceptable. An early process of Castner (B.P. 12218 of 1894), in which molten sodium was allowed to

flow downwards over a column of red-hot charcoal in an iron vessel, while a current of nitrogen was passing upwards through it, did not use the catalytic influence of iron. Castner abandoned it in favour of his synthesis from ammonia and aodium (p. 479), hut it was further investigated by the Roessler and Hasslacher Co in America at a later date. They found that a mixture of sodium vapour and nitrogen reacts with charcoal in the absence of iron above 650°C and that the speed of the reaction above 800°C. is great enough to make the process com-mercially practicable (H. Philipp, Chem. Met. Eng. 1920, 22, 313; U.S.P. 1249821, 1917; 1235897, 1917; 1332439, 1920). Similar proposals are contained in G.P. 368339, 1920. of Broch, who passes nitrogen under pressure through boiling aodium, and heats the vapour and gas under pressure with carbon in presence of an iron catalyst above the melting point of

aodium cyanide. These processes have failed to displace or compete with the established Castner process on account of the great progress which has been

mado in the manufacture of synthetic ammonia.

8. From Carbides and Netrogen - Many proposals have been made for the production of sodium cyanido from carhides and nitrogen They are all either variants of or closely related to the technically important process by which crude calcium cyande is made from ealcium carhide and nitrogen through the intermediate compound calcium evanamide. and are therefore described in the section dealing with this process (Alkalino Earth Cyanides,

p. 487). PROPERTIES OF THE ALEALI METAL CYANIDES Potassium cyanide, KCN, erystallises in anhydrous, colourless octabedra from aqueous solutions, or in cubes from the molten salt The specific gravity of the solid is 1-52. It melts at a dull red heat, and volatilises without decomposition at high temperatures. It is readily solublo in water; the saturated solution at 25°C. contains 71 2 g KCN per 100 g. of water (Bassett and Corbet, J.CS. 1924, 125, 1660), and at the boiling point (103 3°C) 122-2 g, per 100 g, of water. Dry ethyl alcohol dissolves 1-9 g, KCN per litre, methanol 30 per ltre at 25° (Thompson, U.S. Bur. Standards, J. Research, 1931, 6, 1051). It is moderately solubla in anhydrous ammonia; according to Johnson and Krumboltz (Z. physikal. Chem 1933, 167, A, 249), 455 g. of KCN dissolve in 100 g. of ammonia at-33.9°.

deliquesces in moist air. When a mixture of potassium cyanude and and lime is heated in air at 270°C., 23% of the sodal lime is neated in his as 210 0 25 0 true cyanide is oxidised to intrate; silver and nickel assist the oxidisent catigatically, the yields of nitrate being 65 and 71% respectively (J.C.S. 1923, 123, 885) has measured the vapour with these metals. The exidation of eyanide to pressure at various high temperatures: cyanate by alkaline permanganato is accelerated, and secondary reactions are suppressed, by copper sulphato (Gall and Lebmann, Ber. 1928,

thiocyanate, sulphate and thiosulphate; the reaction velocity has been measured by Ishi-kawa (Z. physikal. Chem. 1927, 130, 73), and the effect of varying the pu value of the reacting solution hy Foresti (Atti soc. ital. progresso sci 1934, 22, No. 2, 263; Z. anorg. Chem. 1934 217, 33):

5,0,"+CN'+2OH' #5,0,"+SO,"+CNS'+H.O

This reaction is of physiological importanca in

connection with the use of tetrathionate as an antidote in cyanide poisoning.

A phase rule study of the double salts of potassium cyanide with the eyanides of copper, adver, gold and thallium has been mada by Bassett and Corbet (J.C.S. 1924, 125, 1660). and with those of zinc, cadmium, mercury and nickel by Corbet (sbid. 1926, 129, 3190)

Sodium cyanide, NaCN, erystallises from the molten salt in anhydrous, apparantly cubic creatals. From aqueous solutions below 35°C. large colourless platre of the dihydrate, NaCN,2H₂O, are deposited. Joannis (Ann. chim. phys. 1882, [v], 26, 484) obtained a hydrate NaCN, H₂O from cold 85% alcohol. hydrate NaCN, 1H₂O from cold 85% alcohol. The dihydrato decomposes at 34 7°C. into the anhydrous salt and a saturated solution, about 40% of the sodium cyanide separating in the anhydrous state. The solubility of the dihydrate in water is as follows (in g. NaCN per 100 g. of water):

Temp °C. 34 7 Solubility . 354 43 4 583 The solubility of the anhydrous salt is almost

independent of the temperature at 82 g. per 100 g of water

Dry ethyl alcohol dissolves 97 g. NaCN per litre, and methanol 78 g. per litre at 25° (Thompson, U.S. Bur. Standards, J. Research, 1931, 6 1051). The adubility in aqueous methanol is given below in g. NaCN per litra of

solution: % Methanol by weight. Temp 'C. 100 79 9 80 16 131 207 231 67 9 31 63 3 142

100 g. of anhydrous ammonia dissolve 114 5 g. of sodium eyanide at -12° to -25°C. At lower temperatures a compound NaCN 5NHz separates in large flat plates resembling tha dehydrate. The solid ammoniate is in equilibrium with solutions of the following composition (unpublished work by T. Ewan): Liquid phase, g NaCN

Temp, "C.	per 100 g	
-31	57 7	57-7
-33	43	75
-33	40	80
-34	34	88
-39	21	

Temp °C. 800 900 1000 1100 1200 1300 1350 Vap press (mm.) 076 343 124

beat of evaporation 37 kg.-cals. per g.-mol. The pure salt, when beated in dry air free from carbon dioxide, is stable at temperatures up to 370°C. Rapid oxidation occurs, however, in the presence of iron or nickel or their oxides, with formation of cyanate, carbonate and carbon dioxide. Sodium cyanate is the primary product, which is itself rapidly oxidised to carbonate and carbon dioxide in the presence of nickel:

$2NaCNO+3O=Na_{2}CO_{3}+CO_{2}+N_{2}$

R. Höltje (Z. anorg. Chem. 1933, 214, 65) has investigated the changes occurring in melts of sodium cyanide and sodium hydroxide. In the absence of water and oxygen, reaction only occurs above 500°C., and the products are carbonate, cyanamide and hydrogen. If a little water is present, decomposition starts at 300°C, with saponification to ammonia and formate, the latter being converted into carbonate and hydrogen. Nitrogen, carbonate, and nitrate are produced in the presence of

Sodium cyanide absorbs moisture from the atmosphere. It has been found to take up water from air which has been saturated with aqueous vapour at 20°C. at all temperatures below 44°C. Care is therefore necessary in taking samples of the salt for analysis. Dilute solutions of sodium cyanide slowly decompose in contact with air, the hydrocyanic acid formed in the solution by hydrolysis being lost by evaporation and thus allowing the decomposition to proceed further. Carbon dioxide absorbed from tbe atmosphere also displaces hydrocyanic acid with a similar result. At higher temperatures, for instance, during the evaporation of solutions, saponification to ammonia and formic acid may occur. Data on the rate of bydrolysis of potassium cyanide solutions under various conditions are given by Novalevitsch (Ukrain. Chem. J. 1934, 8, Wiss. tech. Teil, 226) and Wick (Quart. Rev. Amer. Electroplaters' Soc. 1933, 19, No. 10, 20). When strong sulpburic acid is present, the formic acid initially produced by the saponification is immediately decomposed into carbon monoxide and water.

In the presence of oxygen, sodium cyanide dissolves almost all metals with the exception of lead and platinum. In the case of iron, which Liebig found to dissolve in cyanide with production of ferrocyanide and hydrogen, it is necessary for the iron to be in a finely divided form, since iron vessels are used in the technical handling of hot, concentrated cyanide solutions without serious corrosion. Zine is slowly dissolved without oxygen, but more rapidly wben it is present.

Uses of Sodium Cyanide.—The most extensive use of sodium cyanide is in the extraction of gold and silver from their ores, and this subject is dealt with under Gold Cyanides, p. 486. Another use of the alkali metal cyanides is in electroplating, which is mentioned under Silver Cyanide, p. 487. Other applications

are as follows:

(i) Case-hardening.—When a soft steel, usually

The boiling-point is 1,500°C., and the latent | bath of fused sodium cyanide at 800-900°C., both carbon and nitrogen are taken up superficially in the form of a solid solution of the carbide, Fe₃C, and the nitride, Fe₂N, in γ-iron. When the steel is cooled very rapidly this is changed into a supersaturated solution of these substances in a-iron, which is extremely hard; the underlying metal remains unchanged. The composition of the hardened skin may be controlled by diluting the cyanide bath with sodium carbonate, a bath rich in cyanide giving a higher carbon content, and the thickness of the skin by varying the time of immersion; I hour at 900°C., for example, gives a penetration of 0.4 mm. (Beilby and Nelson, J.S.C.I. 1932, 51, 149T). Bucher showed (Ind. Eng. Chem. 1917, 9, 236) that the reversible decomposition of cyanide according to the equation

$2NaCN \rightleftharpoons 2Na+2C+N_2$

takes place in the presence of iron; cyanide and pure iron give carbon and nitrogen by decomposition, which combine with the iron, whereas carburised iron with sodium and nitrogen

gives cyanide.

(ii) Froth Flotation.—In 1922 Sheridan and Griswold (U.S.P. 1427235, 1421585) showed that zinc blende and galena may be separated by carrying out the known froth flotation process in an alkaline solution containing sodium cyanide. Only 0.1-0.2 lb. of sodium cyanide per ton of ore is required to prevent the flotation of the zinc blende, that of the galena being unaffected. The process may also be applied to the separation of the crystalline sulpbides of copper and iron, cyanide acting as a depressant for the iron. A discussion of the mechanism of flotation is given by Brighton, Burgener and Gross (Eng. and Min. J. 1932, 133, No. 5, 276). The action of certain sulphur containing compounds in promoting the collection of minerals in the frotb appears to be connected with their ability to form films of insoluble heavy metal salts at the mineral-solution-air interface, and the action of cyanide in destroying the collecting action is said to be due to the solvent action of cyanide on the collecting films. It is also suggested that, where pyrite is present, the formation of ferroand ferri-cyanides may contribute to the depressing effect; these compounds might become adsorbed at the surfaces, and might also change the polarity of the charge adsorbed on the sulphide surface.

III. HEAVY METAL CYANIDES.

In this section is given a brief account of the preparation and properties of the technically

important heavy metal cyanides.

Copper cyanide.—The only stable cyanide of copper is cuprous cyanide, Cu2(CN)2. When potassium cyanide is added to cupric salt solution, a brownish precipitate of cupric cyanide, $Cu(CN)_2$, is first formed, which gradually evolves cyanogen and changes into green cupric cuprocyanide. On heating, this complex salt loses a further quantity of cyanogen yields white cuprous cyanide, which resembles silver cyanide in constitution and containing about 0.15% carbon, is immersed in a reactions. It may also be produced by the

action of sodium cyanide on a solution of liberated by the solution reaction, by having enprous chloride in sodium chloride solution, present an oxidising agent such as dissolved

when no cyanogen is lost.

Cuprous cyanide is soluble in solutions of the alkalı metal cyanides, about 25 mols. of KCN being required to dissolve 1 mol. of CuCN, the solutions contain the complex amons Cu(CN)," and Cu(CN)," according to Kunschert (Z anorg Chem 1904, 41, 359), but Britton and Dodd (J.C.S. 1935, 100) were only able to find Cu (CN), their conclusion being based on measurements of the HCN vapour pressure over the solution, the p_H value, and the electrical conductivity of the adultion. The complex amon is exceedingly stable, its dissociation into cuprous ions being so slight that no cuprous sulphide is precipitated from the solution by sulphuretted hydrogen. For this reason copper in the presence of oxygen, and all copper compounds, dissolve readily in cyanide solutions. Solutions of sodium and potassium cupro cyanides are used in the electrodeposition of copper on iron.

Gold cyanides .- Aurous cyanide, AuCN, is a yellow crystalline powder, insoluble in water and mineral scids, which is formed by the action of hydrocyanic acid on auric hydroxide, or by warming potassium aurocyanide with hydrochloric seid and evaporating the solution to dryness, washing the residue with water It is readily soluble in solutions of the alkali and alkaline earth cyanides, giving complex aurocyanides such as KAu(CN)₂. A solution of the potassium salt is used for electroplating with gold, and may readily be made by precipitating a solution of gold chloride with ammonia and dissolving the fulminating gold in a solution of potassium cyanide, or by precipitat ing gold chloride with magnesia, dissolving out the excess of magnesia in the precipitate with nitric acid, and then dissolving the auric hydroxide in potassium cyanide as before Sodium aurocyanide is technically important

hecause it is formed when finely divided gold is dissolved in a solution of sodium evanide in the presence of oxygen or other oxidising agent :

4NaCN+2Au+102+H2O =2NaAu(CN)₂+2NaOH

Gold is one of the noblest metals and comes below bydrogen in the electro potential series. Hence when gold is in contact with water or acids, equilibrium in the reaction

2Au+2H+ ≥2Au++H.

is reached at infinitesimal concentrations of simple kation and causes the dissolution reaction | zinc with sodium sulphide : after the addition aurocyanide ion is not in itself sufficient to throwing down a mixture of zine and cuprous necessary, in addition, to remove the hydrogen cyamdation process, and regenerating the zinc

oxygen or hydrogen peroxide. Ionically, there. fore, the reaction by which gold dissolves in acrated evanude solution may be represented as follows.

The dissociation of the complex aurocyanide amon into aurous and cyanide ions is very slight, and smaller than the dissociation of the complex silver and mercury cyanide ions Thus, although free gold is electro negative to both silver and mercury, and would be displaced by them from solutions of simple aurous salts, gold in evanide achitions is electropositive to these metals, and displaces them from their solutions in eyanide. In the case of a more electro positive metal such as zinc, on the other hand, the difference in the solution tendencies of the free metals is too great for the stability of the complex aurocyanide ion to cause a raversal of the normal relation, and zinc readily displaces metallic gold from aurocyanide solutions with the formation of a complex zinc cyanida;

2Au(CN)₂'+Zn⇔Zn(CN)₄"+2Au

In the technical extraction of gold from its ores, the ground ora is leached with alkaline eyanide solution of approximately 0.03% atrength, and the gold is precipitated from the resulting aurocyanide solution by treatment with zinc shavings or finely divided zinc dust. There are frequently present in the ore quantities of iron, copper and zino sulphides which incresso the cyanide consumption in the dissolution process by converting cyanide into complex cyamdes and thiocyanate. In order to reduce the loss of cyamide by this degradation, it is advisable to use cyanide solutions of lower atrength, but a limit to the reduction of strength of the leaching solution is set by the fact that it causes incomplete precipitation of the gold from the pregnant cyanide solution by the zinc current practice at the reduction plants of the

Rand goldfield, the cyanide consumption averages 0 25 lb. NaCN per ton of ora treated. Numerous proposals have been made for the recovery of the cyanide rendered unavailable by reaction with iron, copper, zinc and sulphur in the ore. Mills and Crowe (B.P. 241669, 1924) would ecidify the solution leaving the zine boxes (after precipitation of the gold) with sulphur droxide, and remove the hydrocyanic acid from arrousion. The effect of cyampie, nor and each is re absorbed from the air in shall associate on the effect of cyampie, nor and each is re absorbed from the air in shall except, in which is the each state of the each is real world from the air in shall dissociated complex metal cyanda sons, which the solution and Woolf (Amer. Inst. Min. Met dissociated complex metal cyanda sons, which the solution and then to precupitate copper and the each to the each to the continually reduces the concentration of the continual of the solution and then to precupitate copper and to proceed. In the case of gold, however, the of lime to the filtered solution it is re used, equilibrium concentration of sureas son is R. M. P. Hamilton describes a process whereby a variantingly small that the formation of since sulphate is added to the barren solution, promote the dissolution of the metal, even cyamdes, this is boiled with dilute sulphure though the complex ion, as attack below, is acid, liberating by droyane acid which arternely stable and little dissociated. It is absorbed in milk of lime for return to the

53). In B.P. 416475, 1932, it is proposed to treat the gold-bearing cyanide solution with a cuprous salt, which gives a precipitate eonsisting essentially of cuprous cyanide and the cuprous salts of the complex cyanides of the precious and other metals; this precipitate is treated with sulphuric acid to liberate hydrocyanic acid equivalent to all the complex cyanides, leaving a residue containing the precious metals and cuprous thiocyanate, from which the latter is removed by roasting to copper oxide and leaching with acid. The copper solution may be re-used in the process.

Auric cyanide, Au(CN)3·HCN·1½H2O, is formed on treating potassium auricyanide with

hydrofluosilicic acid.

Mercuric cyanide, Hg(CN)2, is prepared hy dissolving mercuric oxide in excess of hydrocyanic acid and evaporating to crystallisation. It crystallises in anhydrous, colourless crystals of sp.gr. 4.0. It is soluble in water to the extent of 93 g. per litre at 20°C., and 530 g. per litre at 100°C. The solubility in ethyl alcohol is 10·1 g. per 100 g. at 19.5°C., and in methanol 44.2 g. per 100 g. at the same temperature. The electrolytic dissociation of mercuric cyanide in aqueous solution is extremely small, so that mercuric cyanide can be formed, for instance, hy hoiling mercuric oxide with Prussian hlue; this shows that its dissociation is even less than that of the ferrocyanides into cyanide. It is not decomposed by caustic alkalis, and only gives hydrocyanic acid on hoiling with strong acids. Sulphuretted hydrogen does, however, precipitate the extremely insoluble mercuric sulphide from its solution. The reaction of mercurous salts with cyanides yields mercuric cyanide and mercury, instead of the expected mercurous cyanide. On heating to 400°C., mercuric cyanide decomposes into mercury and

Small quantities of mercuric cyanide are sometimes used in addition to alkali cyanide in the treatment of ores containing silver sulphide (Clennel, "Cyanide Handbook," London, 1910, p. 297). By its use the dissolution of silver sulphide is accelerated, probably owing to the formation of mercuric sulphide by the

reaction:

Hg(CN)₂+Ag₂S+2KCN

=HgS+2KAg(CN)2

Silver cyanide, AgCN, is formed by precipitation of silver salt solutions with hydrocyanic acid, or alkali cyanide solution not in excess. It forms a white, cheesy precipitate which is stable to light. Its solubility in water at 25°C. is 4.4×10^{-6} g.-mols. per litre, intermediate between the thiocyanate (less soluble) and the chloride. It is hardly attacked by cold, dilute nitric acid, but is decomposed quantitatively on boiling (Plimmer, J.C.S. 1903, 83, 285). It is slightly soluble in an aqueous solution of hydrocyanic acid, forming the unstable complex acid HAg(CN)₂, and dissolves readily in alkali cyanide solutions to give alkali

sulphate (Eng. and Min. J. 1932, 133, No. 1, sulphide, are dissolved by cyanide solutions; the sulphide and the free metal require the presence also of oxygen for their dissolution (see under Gold Cyanides, p. 486). The soluble salt KAg(CN)₂ is used for electroplating, fulfilling the requirement of producing a tenacious deposit of the proper consistency hy reason of the fact that its solutions have a very small concentration of free Ag+ ions; to ohtain the same conditions with simple silver salts, very dilute and highly resistant solutions would have to be used. The view is held that silver cyanide itself is to be regarded as the silver salt of the acid HAg(CN)2.

On heating in the dry state, silver cyanide gives off about one-half of its cyanide in the form of cyanogen (CN)2, and on further heating in the presence of oxygen gives metallic silver

containing some carbon.

Zinc cyanide, Zn(CN)₂, may be prepared by precipitating zinc acetate solution with hydrocyanic acid vapour. It is also formed by the action of hydrocyanic acid on zinc oxide, but is difficult to prepare in the pure state owing to the ease of its partial decomposition by water. It is a white substance, insoluble in water but soluble in acids and in cyanide solutions. The dry salt decomposes only at a bright red heat.

Complex cyanides, in which the zinc appears in the anion, are formed when zinc cyanide dissolves in alkali cyanide solutions. The complex zinc cyanide ions are much less stable than those containing copper, gold, silver and mercury, and zinc sulphide is precipitated hy a small excess of alkali sulphide. The potassium salt, K₂Zn(CN)₄, crystallises from aqueous solution in large, well-formed octahedra, the solubility of which is 11 g. per 100 g. of water at 20°C. It is fusible at a red heat without decomposition. The sodium salt, NaZn (CN)₃, 2½H₂O, is readily soluble in water.

Barium platinocyanide,

BaPt(CN)4,4H2O,

is made by electrolysing a solution of barium cyanide (270 g. per litre) between platinum electrodes with an alternating current at a current density of 0.4 amperes per sq. cm. About 0.6 g. of platinum is dissolved per ampere-hour. After removing the excess of barium hy means of carbon dioxide, the solution is concentrated to crystallisation (Brochet and Petit, Bull. Soc. chim. 1904, [iii], 31, 630). The salt crystallises in monoclinic prisms, which are yellow by transmitted, and violet-blue by reflected light. It dissolves in 33 parts of water at 16°C. By recrystallisation from a solution containing barum cyanide, it is obtained in small crystals which are highly fluorescent when exposed to X-rays, and are suitable for making fluorescent screens for X-ray work.

IV. ALKALINE EARTH CYANIDES

TECHNICAL METHODS OF PREPARATION .-1. From Carbides and Nitrogen. The direct absorption of nitrogen by the metallic carbides salts of this acid, which are very stable. The leavest of this acid, which are very stable. The leavest of this acid, which are very stable. The light be expected to yield eyanides, and a electrolytic dissociation of the complex $Ag(CN)_2$ considerable amount of work has been done on ion is so small that all silver salts, except the cyanide production. The alkalı metal carbides experiments were carried out up to 1914 in have not been closely investigated, but Frank Germany with httle success. Using 2-3 parts of and Caro (B.P. 15066 of 1895) noted that they absorbed nitrogen at a "dark red heat" to form cyanides. It is possible that the formation of introces was obtained in the form of cyanide from alkali carbonate, carbon and (Landis, Chem. Met. Eng. 1920, 22, 265), mitrogen (see sunder Alkali Metal Cyanides, The technical failure was probably due to the p. 481) may proceed by the primary formation short life of the furnace refrectories, the use of from carbonate and carbon of a carbide, which is unsuitable gas heating, and the occurrence of aubsequently nitrogenated to cyanide, as was frothing in the mass due to the liberation of first suggested by Hughes (Report of Junes, nitrogen.

or mixing and the pure carbides, except that prevented by the addition of small quantities the product from harum carbides except that prevented by the addition of small quantities of ammonia on boiling with vater. In 1897 carbide, The operating temperature was framed to 1.50°C, with marked improvement of Frank and Caro (B.P. 15066 of 1893) discovered the yields (Eastman, U.S.P. 1282305, 1918, that the impure carlides absorb nitrogen, that and the importance of quickly cooling the fused the product in the case of barning carbide reaction mass was realised from the observation

"nnsatisfactory." Similar results were inde-pendently obtained at the same time by Willson crude calcium cyanemide with 0.8 part of salt (B.P. 21997 of 1899), who suggested in the case | and 0.5-0 10 part of calcium cathodus fed into of calcium that a more compound is formed. The fact that a large part type with conducting hearth and a single of the combusting control intogen in introgenated calcium suspended electrods. The matter future, and is carbide (and a smaller part in the case of barnim | run off almost continually into a pan in which sn carbide) is in the form of a cyanamide salt was internally water-cooled drum rotates, taking up

refame and care are accounted to the beautiful of what has been production of cyanides were naturally in marketed have the following analysis Ca(CN), made with barum carbide, which was finely 45 %, NaCl 31-9%, Ca(Cl, 12%, SiO, 93%, Ground and heated with pure intergen to 700°C Fe, Q, and Al, Q, 20%, Ca(CN, 12%, Carbide, Carbi obtained, of which 30% was present as barium cyanide and the rest as barium cyanamide. In order to convert the cyanamide into cyanide, the material was heated with dry sodium carbonate and carbon, when the following reactions may have occurred :

Bancn+ Na,Co, ⇒Baco,+Na,Ncn Na.NCN+C=2NaCN

Proposals to obtain marketable sodium cyanide from this product by extracting the sodium cyanide with water, converting it into ferrocyanide, and then fusing it with sodium by the Erlenmeyer process already described (p. 478)

failed on economic grounds (Erlwein, V. Intern. Kongress angew. Chem., Berlin, 1903, I, 646). Attention was therefore turned to calcum carbide, which absorbs nitrogen at about 1,100°C, giving a product contaming 20-23 6% nitrogen, which is wholly in the form of calcum cyanamide, and about 12% if cathon;

CaC.+N.=CaNCN+C

(For details of the manufacture of calcium cyanamide, see CALCIUM, Vol. II, p. 219). When this product is heated with sodium chlorde, the cyanamide is largely converted and Perret and Perrot (Compt. rend. 1933, 197, into cyanide (Erivein, i.e., Erinem and 761), who studied the reverse of the above Frank, U.S.F. 708333 of 1902). Large-scale reaction, suggest that in the crude cyanife

salt and I part of cyanamide at a maximum temperature of 960°C., from 40 to 75% of the from 40 to 75% of the

The technique of the process was considerably methods as been extensively established. The technique of the process was considerably methods have been extensively established that the process was considerably methods and considerably methods as been extensively established to the process was considerably methods and the process was considerably methods and the process was considerably methods and process was considerably methods and the process was considerably methods. As the process was considerably methods and process was considerably methods and process was considerably methods. The technique of the process was considerably methods and process was considerably methods. consists largely of the corresponding cyanude, that a roversion of cyanide to cyanamide but that the result with calcium carbide was occurs between 400° and 700°C. (Freeman, detouted is in the torm of a yang under Frank's a thin layer of the fused material which coldules direction (Frank and Caro, B.P. 25475 of 1898, and is scraped in in thin flakes. The efficiency paths, Z sugaw, Chem 1903, 15, 6889. Frank and Caro's first attempts at the flakes or powder in which the product is

> The mechanism of this cyanide forming re-action was at first imperfectly understood. The effect of ealt in accelerating the transformation of cyanamide into cyanide is illustrated by the data of W. S. Landis (U.S.P. 1359257, 1920), who heated various mixtures of cyanamide and salt to a temperature above 1,300°C., and then cooled them rapidly to below 400°C., with the following results:

Or	iginal Mixt	ire	Product
Cyanamide.	Salt.	CaC,	%NaCN
200	180	5	36 5
290	100	7	45 0
200	4.5	0	500

Pascal and Bernheim (Compt. rend. 1931, 192, 1425) also found that the yield of cyanide at a given temperature increases with increase in the proportion of salt, though not markedly above a molar ratio NaCl/CaCN, of 2. The optimum proportion of charcoal was 0.2 parts to 1 of cyanamide, which corresponds closely with that actually present in technical crude cyanamide. These results might be explained by writing the fusion reaction as

Canch+C+2NaCl=CaCl2+2NaCN,

obtained by the Landis process the cyanide is combined with sodium rather than with calcium. According to Freeman (Can. P. 229767, 1923), however, the salt/cyanamide ratio in the initial mixture is without influence on the equilibrium yield of cyanide produced in the fusion, and the use of only half the quantity of salt required by the above equation is actually recommended.

recommended.

Landis (U.S.P. 1359257, 1920) had already shown that cyanide formation does occur in the absence of salt, and Franck and Neubner (Z. Elektrochem. 1934, 40, 693) investigated in detail the unassisted reaction between barium cyanamide and carbon, finding increasing cyanide production as the temperature was raised. In the case of calcium cyanamide the relationships are more complicated, and the presence of a flux, preferably a calcium salt such as the chloride or fluoride, is of great advantage. It appears reasonable to assume, with Franck and Burg (ibid. 1934, 40, 686), that the reaction occurring during the fusion is

an endothermic reaction which goes further to the right at higher temperatures (see also Franck and Bank, ibid. 1934, 40, 699). In addition, the slightly endothermic double decomposition,

 $Ca(CN)_2+2NaCl \rightleftharpoons$ $2NaCN+CaCl_2-3.9 \text{ kg.-cal.},$

of which the equilibrium will be little affected by temperature change, may proceed to a small extent. On this view, the crude cyanide obtained from technical cyanamide by fusion with salt consists essentially of a mixture of calcium cyanide and sodium chloride, a conclusion confirmed by an X-ray analysis of the product carried out by Bredig (see Franck et al., Samml. chem. chem.-techn. Vorträge N.F. No. 6, 1931). The conclusion is also in agreement with the general properties of the product, such as its non-hygroscopic character (indicating the absence of calcium chloride) and the rate of its hydrolysis by atmospheric moisture to give hydrocyanic acid.

Crude calcium cyanide is used in gold-mining and in fumigation, but cannot be used in the case hardening of iron and steel. It is at a general disadvantage compared with high-grade fused sodium cyanide on account of its relatively low cyanide strength, and efforts have been made both to convert it into a high-grade product and to medify the synthesis reaction so as to produce a high-grade product directly. Although the desired products are in most cases sodium cyanide, the processes are dealt with at this point, and not under Alkali Metal Cyanides, because of their basic derivation from the Landis

process.

In U.S.P. 1734562, 1929, Cooper proposed to recover sodium oyanide from the crude calcium cyanide by mixing it with sodium carbonate and dissolving the mixture in water to give a solution of sodium oyanide and sodium chloride, which was concentrated to a strength of 20% NaCN, and then cooled to -15°C. to crystallise the dihydrate NaCN,2H₂O. This was dried by

centrifuging, and converted by heating into the anhydrous salt, analysing 80-90% NaCN. This process has not been brought into com-

mercial operation.

An interesting variation of the Landis process is provided by E. J. Pranke, who proposes (U.S.P. 2004130, 1935) to fuse calcium cyanamide with powdered sodium cyanide at 1,350-1,400°C. in an electric furnace; as in the Landis process, the reaction CaNCN+C=Ca(CN)₂ occurs, but in the presence of sodium cyanide a stable double salt Na₂Ca (CN)₄ is formed, which is preserved by rapid cooling. The product as it stands has a cyanide strength equivalent to 80% NaCN and if desired can be converted into pure sodium cyanide by treating with liquid ammonia. Sodium cyanide is dissolved out, and can be recovered by evaporation of the solvent (U.S.P. 1947570, 1934), and calcium cyanide is left, and can be converted into the sodium salt by a method similar to that described in the preceding paragraph (see also B.P. 411177, 1932). In several other patents (e.g. Canad.P 319011, 1932), Pranke claims the production of the double salt from calcium carbide, sodium chloride, and nitrogen, a procedure which constitutes a combination of the cyanamide synthesis and the Landis cyanide fusion reaction in one step, giving a direct nitrogenation of carbide to cyanide. The direct nitrogenation of calcium carbide to cyanide at 1,240°C., with 4% of calcium fluoride as a flux, has been reported by Franck (Samml. chem. chem.-techn. Vorträge N.F. No. 6, 158), who obtained an 85% yield of cyanide (see also G.P. 588943, 1928).

The use of sodium carbonate as a flux instead of sodium chloride is stated by Wöhler, Krall and Dornhöfer (Angew. Chem. 1934, 47, 733) to lead to the conversion of commercial calcium cyanamide into sodium cyanide, the reaction

being:

CaNCN+C+Na₂CO₃=CaCO₃+2NaCN

Whether sodium cyanide is formed during the fusion may be uncertain, but it is obvious that when the product is extracted with water, sodium cyanide solution will be obtained and calcium carbonate left behind. According to B.P. 437614, 1934, fusion of the mass is not necessary, as the reaction proceeds under suitable conditions at 600-1,100°C., and the granular product is then more expeditiously leached with water.

Considerable technical advantages are claimed by D. Hatt (Chim. et Ind. 1932, 28, 777) for a modification of the Landis process which has been investigated semi-technically in France by the Usines de Produits chimiques de Thann. The new process consists in the treatment of calcium cyanamide with a mixture of two substances which react strongly exothermically to give nascent carbon, e.g. calcium carbide and sodium carbonate (G.P. 480905). The production of carbon is said to proceed as follows:

 $Na_2CO_3+2CaC_2=Na_2O+2CaO+5C$ and the whole reaction thus:

CaNCN+2CaC₂+Na₂CO₃=2NaCN+3CaO+4C

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The reaction is exothermic, and starts at 450- eyanamide was transformed into cyanide. An 500°C., after which the temperature rises account of more recent work on the process in spontaneously to 920°C, within 9 minutes. The Sweden is given by Stälhane and Liander crude product contains 20-22% NaCN, and (Ing. Vetenskaps Akad. Hendl. No. 112, 1931. sodium cyanide of 99% purity can readily be produced from it by extraction with liquid ammonia. The crude product of this process is remarkable in that its eyanide content is easily and quantitatively hydrolysed by ateam at atmospherio pressure into acdium fermate and emmonia

So far as is known, none of these modifications to the Landis process has been developed com

mercially. 2. From Carbonates and Nitrogen .- It is evident, from the descriptions already given of methods of fixing nitrogen in the form of alkali and alkaline carth cyanides, that compounds of the alkaline earth metals are more suitable than those of the alkalı metals for fixing atmospherie nitrogen and that, of the alkabne earth metals, the compounds of barrum most readily form cyanides under 'no conditions necessary Margueratte and for nitrogen absorption Margueritte and A. L. de Sourdeval (B P 1171 of 1860, Compt. rend 1860, 50, 1100) discovered that a mixture of barium oxide and carbon readily absorbs nitrogen on heating, and gives barium cyanide. Ludwig Mond (BP 433 of 1882; JS.CI. 1889, 8, 505) also worked on this process, using a mixture of barrum cerbonate and excess of carbon briquetted with pitch eyanido forma-tion was appreciable et 1,200°C, and proceeded most reedily et 1,400°C. The apparatus used was similar to that of Possoz and Boissière in their analogous process with acdium carbonate (see under Allali Metal Cyanides, p. 481), and 40% of the barium was cyanised. In each of the above cases the object of the experiments was the production of ammonia.

As in the technical practice of the Bucher and Landis processes, therefore, internal electrical heating is very auitable. Readman developed 1899 and 1907. Barium carbonate from a later furnace, the lower part of the wall forming one electrods and a central carbon rod suspended in the upper part of the furnace forming the other between the electrodes, and a current of producer upwards through the mass. The cooled product was hyrviated with water, and the solution of barium cyanide cyanamide and hydroxide treated with sodium hicarbonate. After filtering off the barium carbonate precipitate for return to the process, the acdium cyanide solution was only 0 1% carbon monoxide. concentrated in vacuo, cooled to deposit crystals of NaCN,2H₂O which were centrifuged and dehydrated by heating The Badische Amini und Soda Fabrik also developed this process (G.P. Soda Fabrik and overloped time provens (v.r.)
190355, 19040, obtaining a fusion product
containing 60% of its fixed introgen as barum. This decomposition begins at about 500°C,
yearded and 40%, as eyanamide. This was decomposition begins at about 500°C,
and the equilibrium ratios of cyanule/gyanamide
at temperatures from 500° to 900°C, have been

5-60).

A considerable amount of work on the chemistry of this method of nitrogen fixation as eyanide has been done at lower temperatures Bertelsmann (Technologie der Cyanverbind. ungen, 1906, 83) has ascribed the case of nitrogenation of barrum compounds to their infusi hility, so that they remain solid and present a large aurface to the action of nitrogen under the eyanising conditions. But although unde composed barium carbonate does not melt at 1,350°C. (Finkelstein, Ber. 1906, 39, 1585), its mixtures with harium oxide, which are eluque formed at the high temperatures in question, are readily fusible. The formation cyanide (m.p. about 600°C.) still further increases the fusibility of the reaction mixture, which must therefore in this process be a liquid phase of variable composition.

Kuhing and Berkhold (Ber. 1908, 41, 28) obtained 40% nitrogenation of the barum in a mixture of harium carbonate and carbon at 1,150°C., the reaction diminishing as the temperature was lowered and being extinguished at about 925°C Ewan and Napier (JSCI 1913, 32, 467) showed that nitregen absorption which, as stated by Mond (I c.) depends on the two reversible reactions,

 $BaCO_*+C \rightleftharpoons BaO+2CO-1012 \text{ kg.cal}$ $BaO+3C+N_2 \rightleftharpoons Ba(CN)_2+CO-43 l kg cal$

is therefore hindered by the accumulation of carbon monoxide, and favoured by the use of a large excess of nitrogen. In 2 hours at 960°C, they obtained 40% nitrogenation of the benum when using a 28-fold excess of nitrogen. The pressure of carbon monoxide in equilibrium with pure herium carbonato and carbon is large enough to prevent the second reaction from such a process (BP, 6621 of 1894), which was proceeding to the right, but as the banum worked by the Scottish Cyanides Co. between carbonate is diluted with barium oxide the equilibrium pressure falls to very low values stage was mixed to a thick pasto with ground It is therefore concluded that the greater part coal, dried and coked in ordinary gas retorts of the barum carbonate decomposes before any The granular mixture of barum oxido and coke of evanido is formed. Evan and Napier obtained ao produced passed to a cylindrical electric the same equilibrium product, at a given tem perature and gas composition, from starting mixtures of BaCO3+BaO+C and of Ba(CN)+ BaCN4+C. Their results at temperatures of The solid mass was heated in passing downwards 1,000°, 1,100°, and 1,150°C, with vanous mixtures of nitrogen and carbon monoxide gas containing 70% of nitrogen was passed appeared to indicate by extrapolation that the maximum quantity of barium which could combine with pure nitrogen was 50% at all the temperatures, but they obtained a product corresponding to 57 1% nitrogenation by heating for 82 hours at 1,100°C, with nitrogen containing

> A further equilibrium which must be taken into account is the decomposition of cyanide into eyanamide and carbon:

measured by Franck and Neubner (Z. Elektrochem. 1934, 40, 693), who find that only 3% decomposition of cyanide occurs at 900°C. According to Ewan and Napier, the presence of barium oxide in the melt has a disturbing effect on the equilibrium, so that the relative amounts of cyanide and cyanamide produced by the action of nitrogen on mixtures of barium oxide and carbon are very erratic. They do not consider that barium carbide can be an intermediate product, since the pressure of carbon monoxide present is always greater than the equilibrium partial pressure which they determined for the reaction

$BaO+3C \rightleftharpoons BaC_2+CO.$

Hempel (Ber. 1890, 23, 3388) and Stähler (Ber. 1916, 49, 2292) found that the formation of cyanide was greatly favoured by working under a nitrogen pressure of 60 atmospheres, Stähler obtaining a product containing 74% of barium cyanide with no cyanamide. Caro of barium cyanide with no cyanamide. Caro (Z. angew. Chem. 1910, 23, 2405) studied the effect of adding calcium and barium fluorides to the melt at 1,100°C, and 1,200°C. In no case was the barium nitrogenated to a greater

proportion than 50%.

At higher temperatures it appears that as much as 80% or more of the barium can be nitrogenated (Badische Anilin und Soda Fabrik, B.P. 2133 of 1907, 22039 of 1906; Kaiser, F.P. 454237, 1912; 454238, 1913). Ewan and Napier suggest that, at temperatures above 1,200°C. barium carbide may be an intermediate product, since by applying Nernst's theorem to their measurement of the carbide-forming reaction mentioned above, they were able to calculate that the equilibrium pressure of carbon monoxide at 1,500°C. is 100 mm., and at 1,700°C. is 1,026 mm. These figures suggest that the carbide could exist under the experimental conditions obtaining at these temperatures. In this connection it may be noted that Franck and Neubner (Z. Elektrochem. 1934, 40, 693) noticed the formation of carbide, as well as cyanamide, on heating barium cyanide above 900°C.

3. From Hydrocyanic Acid.—It is difficult to preparc pure anhydrous barium cyanide from the aqueous solution which is obtained by neutralis-ing baryta with hydrocyanic acid, owing to the ready hydrolysis of the salt. In the case of calcium cyanide, which is a technically valuable compound, the pure anhydrous salt can under no condition be prepared from aqueous solutions, the hydrolysis being assisted by the formation of the relatively insoluble calcium hydroxide. An impure technical anhydrous calcium cyanide may be made, however, by the interaction of calcium oxide and hydrocyanic acid, both in the anhydrous state. The Deuts. Gold-u.-Silber-Scheideanstalt describes, in G.P. 555223, 1929, the preparation of a powdered product containing more than 40% Ca(CN)2 by treating powdered calcium oxide with boiling anhydrous HCN. The presence of 1-3% (by weight of the HCN) of an accelerator such as ammonia or water is recommended in order to minimise the loss of HCN by polymerisation. The reaction may also be effected in an inert medium such as diethyl other, in which calcium cyanide is insoluble. Similar proposals have been made by the Roessler and Hasslacher Co. (U.S.P. 1909042, 1933; 1950879, 1934).

Franck and Freitag (Z. angew. Chem. 1926, 39, 1430) describe a method of making calcium cyanide of 99% purity by the action of hydrocyanic acid on a solution of calcium in liquid ammonia, when hydrogen is evolved and a precipitate of the diammoniate,

Ca(CN)2,2NH3,

is produced. The same compound may also be obtained by the action of HCN on a solution of calcium nitrate in liquid ammonia. It is stable in vacuo at 100°C., but begins to lose ammonia at 150-160°C., and is completely free from it after 2 hours at 180°C. Similar data are given by Poindexter in U.S.P. 1596120, 1926; 1652874. 1927. A manufacturing on these lines has been developed by the I.G. Farbenindustrie A.-G., and is described in B.P. 300349, 1927. A calcium compound such as the oxide or nitrate reacts with hydrocyanic acid and ammonia in the presence of water to give small crystals of the diammoniate, which are dried by washing with an organic liquid taking up water and are then decomposed by heating to give a product of the following analysis: Ca(CN)₂ 85-88%, Ca(OH)₂ 8-11%, CaCO₃ 1.0%, NH₃ 0.5%, which is used in fumigation.

Another method of manufacture of calcium cyanide is described in Metzger's B.P. 261722, 1925, and in Ind. Eng. Chem. 1926, 18, 161. Liquid hydrocyanic acid reacts with calcium carbide according to the equation

$$CaC_2+2HCN=Ca(CN)_2+C_2H_2$$
.

The reaction is carried out by stirring powdered carbide with excess of the liquid to complete the reaction, distilling off the excess and leaving a finely divided buff powder of the approximate composition Ca(CN)2,2HCN, from which the hydrocyanic acid may be easily removed by heating in vacuo. According to Franck and Freitag (l.c.), the product from commercial carbide contains not more than 55% Ca(CN), with 25% HCN, and after removing the hydrocyanic acid leaves impure calcium cyanide containing 25-30% CN.

Properties of the Alkaline Earth Cyan-IDES.—Barium cyanide. Crystals of the dihydrate, Ba(CN)₂,2H₂O, may be prepared by adding hydrocyanic acid to baryta in tbeoretical quantities, and evaporating the concentrated solution in vacuo (Joannis, Ann. chim phys. 1882, [5], 26, 484). The evaporation may be avoided by drying pure Ba(OH)₂,8H₂O, in vacuo at 100°C., and suspending the finely divided powder of anhydrous barium hydroxide in dry light petroleum to be treated with a small excess of dry hydrocyanic acid also mixed with light petroleum; the dihydrate then crystallises directly:

$Ba(OH)_2+2HCN=Ba(CN)_9,2H_9O$

The dihydrate is very deliquescent, but may be converted into the amorphous monohydrate by drying in vacuo over sulphuric acid, and the 492 CYANIDES.

racuo. Although barnum cyonide is much more readily soluble in weter but not in hydrocyanic atable than the other alkaline earth cyanides, acid, is precipitated and washed with hydroit is difficult to evoid some hydrolysis during the cyanic acid and dried in vacuo at 40°C. for at above dehydration, so that the product often least 30 minutes. B P. 314242, 1928, recom contains some barium hydroxide, which can, however, be removed by shaking the solid with a small quantity of bydrocyanic acid and petroleum and redrying, giving a practically pure product. If methanol is used instead of petroleum in the shove preparation, a com pound, Ba(CN)(OCH₂)+CH₂OH 28 obtained, which on heating loses first methanol sacuo above 250°C. and then dimethyl ether, leaving

BaO+Ba(CN).

(Drechsel, J. pr. Chem, 1880, [11], 21, 77)

Dry methods of preparation always yield mixtures of cyanido and cyanamide; for instance, by heating harrom ferrocyanide above 500°C, or by the ection of nitrogen on barium carbide the same product, containing about 60% of its nitrogen as cyanamide, is obtained This is due to the equilibrium

Ba(CN), ≠ BaNCN+C,

which has been studied, as already mentioned, hy Ewen and Napier (JSC.I 1913, 32, 467) and by Franck and Neubner (Z. Elektrochem 1934, 40, 693).

Barum cyanide is a colourless solid, melting at 500°C. opproximately, at which temperature it is appreciably volatile. It is soluble in water to the extent of 80 g. per 100 g. of water at 14°C, and much less soluble in alcohol.

Calcium cyanide cannot he prepared in the wet way. An aqueous solution of lime in hydrocyenic acid decomposes almost completely on bolling, and on evaporation in tacuo gives a crystalline compound, 3CaO Ca(CN), 15H, O (Joannis), which decomposes completely in a vecnum, leaving only lime. Dry reactions at high temperatures, on the other hand, yield only calcium cyanamide, for in the case of the calcium compounds the cyanide-cyenamide equilibrium is more in favour of the cyanamide,

Finely powdered calcium cyanide is rapidly hydrolyaed by atmospheric meisture, with almost quantitative evolution of hydrocyenic avoid accondary decompositions acid;

On account of this property it finds use as a

Magnesium cyanide has not been prepared, but is known in the form of a diammoniate corresponding to the calcium compound. This ammonia with magnesium. Olberg obtained it

monohydrate in turn may be completely excess of hauld hydrocyanic acid (UEP, debydrated by gradual heating to 100°C. on 1609038, 1926). The compound, which we mends the use of ethanol instead of water in the preparation, since the compound is practically msoluble in this solvent end the use of a large

excess of hydrocyanic acid is therefore avoided.

Mg(CN), 2NH, is more stable than the corresponding calcium compound, the ammonia being removed only by prolonged heating in

V. HYDROCYANIC ACID.

TECHNICAL METHODS OF PREPARATION -1. From Ferrocyanides.—Ferrocyanides give bydrocyanio acid on treating with dilute sulphune seid (Gattermann, Annalen, 1907, 357, 318). Only one-half of the cyanide is thus recovered as HCN, however, owing to the formation of the relatively insoluble Eventte

 $2K_{4}Fe(CN)_{4}+3H_{5}SO_{4}$ =6HCN+ $K_{2}Fe_{2}(CN)_{4}+3K_{5}SO_{4}$

Everitt's ealt can be incompletely reconverted anto soluble ferrocyanide by houling with caustro soda (Erlenmeyer), the complete con version requiring the assistance of simultaneous oxidation with air, when Fe O4 or ferror hydroxide is precipitated instead of ferrors hydroxide (Grossmann, B.P. 36, cf. 4513, both of 1903). Dewrance and Williams (B.P. 28074 of 1908) ohtain complete decomposition of alkali ferrocyanides by distilling them with acid in the presence of a small amount of cuprous chlonder the action is supposed to be due to the intermediate formation of cuprous ferrocyands having the property, exceptional among ferrocyenides, of yielding the whole of its cyanogen as hydrocyanic acid on decomposition with boding dilute sulphuric acid. The decomposition of complex iron cyanides with steam in the absence of ands is mentioned in F.P. 721762, 1931; rapid removal of the hydrocyanic and from the reaction zone is necessary in order to

2. From Thiocyanales, -The reduction of alkali thiocyanates with hydrogen, when e part of the cyanide produced appears as hydrocyans acid, has already been mentioned. With barum Transgard, which is applied to the space to a sea and emporate the funigated by blowing the fine powder as a Sones, J.S.C.I. 1901, 20, 320) the pranciple cloud with an frome band-operated fan Thydrolytic decomposition is completed in a few the case of the cuprous all, the readies minutes. Albright (B.P. 4403 and 6226 of 1901) mixed cuprous thiocyanete with finely divided copper and heated the mixture with hydrogen at compound was first obtained by Bergstrom temperatures rising from 150° to 550°C. Hydro (J. Amer. Chem. Soc. 1924, 48, 1565) by cyante acid was evolved and cuprous sulphide treating a solution of mercurar cyanules in hagud cyanic acid was evolved and cuprous sulphide

The most important method of converting hy treating a saturated aqueous solution of thiocyanetes into hydrocyanic acid is Raschen's magnesium nitrate or chlorude (1 mol.) with mitric acid oxidation process (B.P. 10476, 10956, 4-6 mols, of ammonia, and then with a large and 21678 of 1895; 19767 of 1898; 12180 of CYANIDES. 493

1900), which has been operated on a commercial scale (Conroy, J.S.C.I. 1899, 18, 432). A 15% sodium thiocyanate solution was allowed to flow into dilute nitric acid kept at the hoiling-point by means of live steam:

NaCNS+2HNO₃=HCN+NaHSO₄+2NO

A slight excess of nitric acid was used in order to obtain a quantitative decomposition. The gases leaving the decomposer were scruhhed with a limited amount of water to remove traces of N₂O₃, the wash water heing returned to the decomposer. The 33% of HCN in the resulting gas was actually removed hy absorption in caustic soda solution of sp.gr. 1.37, avoiding the presence of an excess of hydrocyanic acid owing to its tendency to form polymerisation products in the presence of cyanide, and the cyanide solution evaporated to dryness in vacuo. A 96-99% yield of HCN is claimed. The nitric oxide remaining was mixed with air and the mixed gas scrubbed in flint-packed towers with water, giving a nitric acid solution of sp.gr. 1.12 which was returned to the system. Two more recent patents of the Gesellschaft für Kohlentechnik (B.P. 356190 and 356724, 1929) describe a similar process starting ammonium thiocyanate, a readily available by-

product from gas works.

3. By the "Schlempe" Process.—The juice of the sugar heet contains both inorganic and organic substances other than sugar, and these substances accumulate in the molasses. In Germany, the greater part of the molasses produced is treated with strontia, which combines with the sugar to form an insoluble sucrate. The liquid residue from this precipitate, which contains the impurities of the original juice in concentrated form, is known as "schlempe," and is concentrated to a sp.gr. of 1.4. It then contains 75% of dry substances, of which 30% are inorganic (mainly potassium) salts and the rest organio substances. It contains about 4% of nitrogen in the form of hetaine,

and decomposition products of the vegetable proteins. A small part of the molasses which is not treated by the strontia process is fermented to alcohol, and the residue from this fermentation is known in France as "vinasses." When "vinasses" are distilled at 700-800°C. in horizontal iron retorts, vapours containing ammonia, the three methylamines, various higher alkylamines, methanol and combustible gases are evolved, and a solid mixture of potassium carhonate and carhon remains hehind (Duvillier and Buisine, Ann. chim. phys. 1881, [5], 23, 289; Ost, Z. angew. Chem. 1906, 19, 609). Vincent worked this distillation process in 1877–1880 for the preparation of methylamines, and in 1879 Ortlieh and Müller developed a process for converting the methylamines into hydrocyanic acid, ammonia, methane and hydrogen hy passing their vapours through a red-hot retort. This process was operated by the Société Anonyme de Croix

(B.P. 3844 of 1879), the ammonia being absorbed in sulphuric acid and the hydrocyanic acid converted into ferrocyanide by treatment with a suspension of ferrous hydroxide in caustic potash. The manufacture of methylamines was abandoned in 1881 for want of a market, and with it the subsidiary production of cyanides.

The production of hydrocyanio acid from "schlempe" was put on to an economic hasis hy Bueb (see Reichardt and Bueb, B.P. 7171 of 1895; Bueb, B.P. 26259 of 1898), who cut out the intermediate separation of the methylamines and passed the "schlempe" distillation products directly to the cyanising furnace. Bueh's process (Muhlert, Chem. App. 1925, 12, 156) is worked at the present time as follows: the liquid "schlempe" is run into a horizontal retort of the gas works type made of highly aluminous fireclay and heated by the residual gas formed in the process. The pasty carbonaceous residue has to be scraped out of the retort hy hand, and is then lixiviated to recover potassium carbonate. The gases, after condensation of tar in a hydraulic main, pass directly to the cyanising chamhers, which are filled with fire-brick checker-work. During the cyanising, which is carried out at above 1,000°C., carbon is deposited on the brickwork and the temperature falls; reheating is carried out by direct firing with producer gas, when the carbon deposit is burnt off. Two such chambers are therefore used alternately, one being reheated whilst the gases are being cyanised in the other. The cyanised gas contains 7% HCN, 7% NH₃, 8% hydrocarbons (mainly methane), 12% H₂, 18% CO, 24% CO₂, and 24% N₂. It is cooled and scrubbed with hot dilute sulphuric acid to remove ammonia. The hydrocyanic acid is then removed by water scrubbing in a 24-plate bubble cap column, a 2-3% aqueous HCN solution being obtained. This is distilled through a fractionating column having 11 plates. The hydrocyanic acid leaves the top of this column as a concentrated gas.

In the early days of the process, the hydrocyanic acid was worked up into ferrocyanide by the wet process for HCN-recovery from coal gas (see Ferrocyanides, p. 468). At the present time, however, it is converted into relatively pure hriquetted sodium cyanide hy the method described under Alkali Metal Cyanides, p. 481. Of the nitrogen in the "schlempe," about 45% is recovered as sodium cyanide, 20% as ammonium sulphate, and 35% is lost as elementary nitrogen.

The decomposition of monomethylamine at a red heat into hydrocyanic acid, ammonia, methane and hydrogen was noted by Wurtz (Ann. chim. phys. 1850, [iii], 30, 443), who

established the following equation:

$3CH_3NH_2=2HCN+NH_3+CH_4+3H_2$

process in 1877–1880 for the preparation of methylamines, and in 1879 Ortlieh and Müller developed a process for converting the methylamines into hydrocyanic acid, ammonia, methane and hydrogen hy passing their vapours through a red-hot retort. This process was operated by the Société Anonyme de Croix The hehaviour of di- and tri-methylamines under similar conditions, when passed through heated porcelain tubes containing broken porcelain, firebrick or charcoal, has heen studied by Voerkelius (Dissertation, Hannover, 1909). The formation of hydrocyanic acid from dimethyloperated by the Société Anonyme de Croix

to the equation

occurs. Increasing dilution of the dimethylsmine vapour with hydrogen leads to the production of ammonia by the reaction

and 2% according to

$$NH(CH_3)_2 + 2H_2 = NH_3 + 2CH_4$$

With a gas containing 50% hydrogen hardly any emmonia is formed, but with 97% hydrogen no hydrocyanic acid is produced results were obtained with trimethylamine, which at temperatures between 800° and 1,000°C decomposes to the extent of 98% according to the equation

N(CH₃)₃=HCN+2CH₄,

 $N(CH_3)_3 + 3H_3 = NH_3 + 3CH_4$

Within the above temperature range, the results are not much affected by changes in the time of contact or the amount of heated surface exposed to the gas But at higher temperatures, or in the presence of iron or refractory materials at longer times of contact, considerable decomposition of the hydrocyanic acid into its elements occurs At 600°C no decomposition of the trimethylemine occurred to give hydrocyanic acid, end at 720°C the yield was only 78%

The Deuts Gold u. Silber Scheideanstalt vorm Roessler (F P. 447725, 1911, and 448722, 1912) clum that, by passing the "schlempe" gazes at high velocity through heated passages made of fused quartz or of Dinas brick without filling material (thus heving the smallest possible con tact surface), the whole of the nitrogen is obtained as ammonia or hydrocyanic acid According to GP 420729, 1924, the salts of trimethylamine, betains end hexamethylenetetramine (especially the hydrochlorides) give hydrocyanic acid on heating in greater yield than is obtained from the free bases, betaine

quantity of HCN as compared with 50-60% obtainable from betains itself 4 From Ammonia and Carbon Monoxide -The production of hydrocyanic acid from carbon monoxide and ammonia by heating them in the presence of spongy platinum was first noticed by Kuhlmann (Annalen, 1841, 38, 62) Jackson and Laurie (J.C.S. 1905, 87, 433), working with the dry gases, were unable to confirm this observation, their only products on sparking, treating by the silent electric discharge, or in the

hydrochloride giving 70-80% of the theoretical

presence of a heated platinum wire being ammonium eyanate and bydrogen . CO+2NH,=NH,CNO+H.

Mailhe and de Godon (Bull Soc chim. 1920, [1v], 27, 737) found that alumina catalysed the production of hydrocyanie acid at 400-420°C. a rapid reaction occurring at 550°C

Many catalysts have since been proposed in the CaSO₄+2NH₃+CO₃+H₄O

—CaCO₃+(NH₄)₅SO₆

1,100°C. quantitative decomposition according thorsa and alumina (Badische Andin und Soda thora and atumna (baussine atum und cons Fabrik, B.P. 220771, 224438, 1923; IG. Far bennid, A.-G., B.P. 200369; Dents, Goldo, Silber Scheidenstalt vorm. Roessler, B.P. 207830, 223918, 1922; Bredig and Elöd, BP 229774, 229973, 1923; G.P. 522532; USP. I598707, 1926; 1634735, 1927; Frank and Cam, BP. 282379, 1926; 301093, 1927; CP

611726). Further information on the catalytic reaction is given in the technical literature by Bredic and Elod (Z Elektrochem. 1930, 36, 1003; 1931, 37, 2), who worked with alumina, ceris and thoras at 500°, 600°, and 700°C and at reaction times of 1-12 seconds With a gas containing 20 volumes of carbon monoxide and I of ammonia, a contact time of I second at 700°C with alumina gave a 65% yield of hydracyanic acid from the ammonia, and 140, of the ammonia was lost by decomposition; with a volume ratio of 10CO.1NH, under the same conditions, the HCN yield was 48% Fuchs and Verbeck (Ind Eng. Chem 1935, 27, 410) describe the production of an active alumina catalyst, probebly in the form of a gel, by precipitation of aluminium hydroxide from aluminium salt solutions and drying the pre cipitate under controlled conditions catalyst at 596°C, a gas mixture of inital composition 9CO 1NH, gave a 54% conversion of ammonia into hydrocyenic acid with a contact time of I second

Yields of this magnitude are surprising in view of the case with which the reverse reaction. the hydrolysis of hydrocyanic acid to ammonia and carbon monovide, can be carried to com pletion by heating with steam and a catalyst euch as broken brick, iron oxide or manganese dioxide (see Carpenter and Linder, JSCI 1905, 24, 63). The explanation is probably to be found in the reduction of the partial pressure of steam by the operation of the water gas equilibrium.

.CO+H.O ⇔ CO.+H.,

for which the equilibrium constant

[CO][H,O]/[CO,][H,]

at 600°C, as 0 35 Bredge and Elod (Z, Elektro chem. 1930, 36, 1003) find that the use of an excess of carbon monoxide produces a greater merease in the HCN yield than a corresponding excess of ammonia, which is in barmony with the above explanation. The reaction mechanism when ceria is used as the catalyst is discussed in

Z Elektrochem 1931, 37, 2

The separation of the hydrocyanic acid from the reaction inixtures offers difficulties, since unchanged ammonia and carbon dioxide are present in the gas Many of the methods described under Alkalı Metal Cyanides, p 480, are applicable, and in addition it has been proposed (1) to freeze out ammonium cyande (G.P. 444504, 1922), (2) to treat the ammonia and carbon dioxide with calcium sulphate and

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of ammonium sulphate (U.S.P. 1606767), and (3) to absorb the hydrocyanic acid in active charcoal or silica gel. An interesting adaptation of the reaction to the direct production of fused sodium cyanide has been proposed in B.P. 301565, 1927, and in U.S.P. 1955229, 1934; a mixture of ammonia and carbon monoxide is caused to react with sodium hydroxide, carbonate, sulphide or other salt at 580-650°C.

5. From Formamide.—Hydrocyanic acid can be obtained by the catalytic dehydration of

formamide:

H-CO-NH2=HCN+H2O

Formamide itself may be prepared by the

following methods:

(a) From formic acid and ammonia, which give ammonium formate. On distilling ammonium formate, water containing small quantities of ammonia and formic acid is first given off, and at 180°C. the distillation of formamide begins. As the temperature is raised to 220°C., increasing quantities of carbon monoxide are evolved. The reactions occurring are:

$$\begin{array}{l} \text{H} \cdot \text{CO} \cdot \text{ONH}_4 \rightarrow \text{H} \cdot \text{CO} \cdot \text{NH}_2 + \text{H}_2 \text{O} \\ \text{H} \cdot \text{CO} \cdot \text{ONH}_4 \rightarrow \text{H} \cdot \text{CO} \cdot \text{OH} + \text{NH}_3 \\ \rightarrow \text{H}_2 \text{O} + \text{CO} + \text{NH}_3 \end{array}$$

Experiments on this decomposition are described by Lorin (Annalen, 1864, 132, 255) and by Freer and Sherman (Amer. Chem. J. 1898, 20, 223), who recommend that the dehydration be carried out in an atmosphere of ammonia. The use of dehydrating catalysts such as bauxite for the vapour phase reaction at 170°C. is protected by the Badische Anilin und Soda Fabrik in B.P. 237528, 1925).

(b) From alkyl formates and ammonia, the reaction having been discovered by A. W. Hofmann (J.C.S. 1863, 16, 72), who saturated ethyl formate with ammonia and heated it in a sealed tube at 100°C. for 2 days. Alkyl formates are produced by treating solutions of sodium alkoxides in the corresponding alcohols with carbon monoxide under pressure (Stähler, Ber.

1914, 47, 580):

CO+NaOMe=Na·CO·OMe HOMe+Na·CO·OMe

≈H·CO·OMe+NaOMe

At the ordinary temperatures the reaction is slow, but goes to complete ester formation on prolonged treatment with carbon monoxide at 300 atmospheres; at 190°C., however, only 4% of the alkoxide reacts. Similar results were obtained by Fischer and Tropsch (Ges. Abh. Kennt. Kohle, 1921, 6, 382), using sodium formate as catalyst, and by Christiansen (J.C.S. 1926, 129, 418) on passing a mixture of carbon monoxide and methanol vapour over sodium methoxide at 70-100°C., high pressures and low temperatures favouring the formation of ester. According to B.P. 252848, 1925, when methanol containing 10 g. of dissolved sodium per litre is heated to 80°C. with carbon monoxide at 60 atmospheres, 75% of the alcohol is converted into methyl formate in 3 hours, whilst at 1,000 atmospheres the reaction is complete in a few minutes.

as in a technical method for the manufacture | reaction is described in B.P. 240087, 1925. Methyl formate is first made by dissolving sodium in methanol and treating the solution with carbon monoxide as above. The product is then treated with ammonia at 60°C. and 150 atmospheres. B.P. 254787, 1925, describes a vapour phase method by which methyl formate vapour is passed with ammonia over a dehydrating catalyst such as alumina, thoria or silica gel at 200-300°C, with a short time of contact: at higher temperatures or with longer contact times the principal product is hydrocyanic acid.

(c) From ammonia and carbon monoxide directly. K. H. Meyer and Orthner (Ber. 1921, 54, [B], 1705) obtained some formamide, together with ammonium carbonate, formate, and cyanide, on heating a mixture of carbon monoxide and ammonia at 200°C. and 230 atmospheres in the presence of earthenware. A similar process is described in B.P. 203812, 1922, but is not of technical interest owing to the predominance of side reactions. The process described in B.P. 240087 above may be carried out by introducing the ammonia and carbon monoxide simultaneously into the sodiummethanol solution under the stated conditions, when it would fall under this head. The R. and H. Chem. Co. describe, in U.S.P. 1787483, a similar reaction at e.g. 70°C. and 17 atm.

The dehydration of formamido to yield hydrocyanic acid will obviously require temperatures higher than 200°C., since formamide itself is produced by dehydration of ammonium formate at that temperature. From thermochemical data it may be calculated that the reaction is endothermic, the equilibrium therefore favouring HCN-formation at high temperatures, and it proceeds with an increase in gas volume, so that the yields of hydrocyanic acid are greater at lower pressures. The variation of the equilibrium

constant $k_p \left(= \frac{[HCN][H_2O]}{[H \cdot CO \cdot NH_2]} \right)$ with temperature is given by the relation

$$\log k_p = -\frac{23380}{4.571\text{T}} + 1.75 \log \text{T} - \frac{0.0046}{4.571} \text{T} + 3.5,$$

the partial pressures of the components being expressed in atmospheres. From this relation, the percentage conversion of formamide to hydrocyanic acid at equilibrium at various temperatures and under total pressures of l and 0.02 atmospheres may be calculated as follows:

80	Percentage conversion.		
Temp. °C.	1 atm.	0 02 atm.	
200 300 350 400 500 600	8·8 25·0 	53·1 87·7 97·3 99·4 —	

Under favourable conditions the yields obtained The manufacture of formamide by this in practice approximate to those calculated.

CYANIDES.

In the technical practice of this theoretically rery favourable reaction for the production of hydrocyanic acid, several difficulties arise. It is found that liquid formanide near its boiling-point decomposes rapidly, hough the vapour is relatively stable. For this reason it is accessive that the liquid be help at its boiling necessary that the liquid be help at its boiling mannonia, and of the modifications of and in provenance in the liquid be help at its boiling hearing the method of cyaning amonia, and of the modifications of and in provenance in the simple reactions which have received in the modification of and in provenance in the simple reactions which have received in the modification of the modification of and in provenance in the simple reactions which have received in the modification of the modificatio isguid (preferably under reduced pressure) on to a superheated surface kept considerably above 215°C, so that vaporesation is immediate and no loqued accumulates in the oraporate (R.P. 301974, 1927). Side reactions may 1916, pyridine and its homologues (R.P. 331924, 1927). Side reactions may 1916, pyridine and its homologues (R.P. tant being the reaction of hydrocyanic acid with each steam to give carbon monoxide and ammond. The use of catalysts which accelerate formamde challenges that the brightness of the catalysts at such as according to the citylamuse (R.P. 33193, 1927). When metallic catalysts at steam to give carbon monoxide and authon the superhatures are used, for example the catalysts which accelerate formamde copier in the form of a wire net at 487°C. dehydration but which do not favour the secondary decomposition as therefore advantage. The most important catalysts or lattisty, 1915. Bredge, Eliod and Dennas (April 1915) Bredge, Elio in promoting HGN-accomposition (B F 300816, acceptions and ammonia at 600°C, and central 1928), and certain metals such as harsa and times of 1.4 seconds through found 29% of the astel (U.S.P 1678366, 1928, and 1951520, 1934), ammonia converted into hydrograma and pressure (B.P. 260166, 1928, and 261857, 1925), of the actylene was documposed into asthe or or with formamide vapour diluted with ammonia (B.P. 233080, 1924), is also mentioned

The last technical problem presented by this process is the supply of the heat quantities necessery to maintain the endothermic reaction C₂H₈+2NH₈=2HCN+3H₂. The use of an at the high space velocities employed. Heat alumina estalyst for the formation of hydroman space vectories employed. HeAN aumman estalyst for the formation of hydrotransfer is probably aided by the use of metalloc earns are from ammona and enter-setyless tubes filled with metallic catalyst as reaction or henzens is mentioned in G.P. 55505, 1827.

(a) Englage and Ammona also react to give 285200, 1928, describes a reaction chamber having brass walls heated to a temperoture above 300°C, no portion of the formamide being that precepitated alumina mixed with quart more than helf an inch from the heated walls was the best catalyst, which gave the following A further method of ohtaining rapid heat results at 800°C, with a gas mixture consisting of supply is to pass the vapour of formamide equal volumes of ethylene and ammonie: through a bath of fused metal heated to 300-600°C., GP. 561816, 1930, clarms 70-85% yields of hydrocyanic acid by this method Finelly, it is claimed in BP 292749, 1927, that an alumina cetalyst may be deposited on a wire gauzo which can be heated electrically during the passage through it of formamide vepour.

6. From Ammonia and Hydrocarbons .- Hydro cyenic acid is produced by a large variety of gaseous reactions between ammonia and carbon compounds. Since the synthesis of HCN from its elements is highly endothermic, these reactions are generally favoured by high temperatures, and the reaction temperature Considerable quantities of methane and carbon required for HCN formation rises in passing were also formed. The same catalyst was from the most unsaturated hydrocarbon, found effective by Eled and Nedelmann (Z. acetylene, through ethylene to the completely Elektrochem. 1927, 33, 217) for the reaction saturated methane. With the unsaturated between ethyline and nitrio oxid at 80g-hydrocarbons, eatalysts may usefully be 950°C. Probally the nitrio oxide as first reduced comployed to accelerate the oyemastine at to amonia, which their reacts with the temperatures between 500° and 900°C, but with decomposition methano very high temperatures are required (C.H.=CH.+CH.; and CH.+NH (above 1,000°C.), and very short times of contact —HCN+2H.). Using a gas mixture con nust be used in order to minimise loss by taining INO 2C₂H₄ 2H₄ at a temperature of secondary decompositions. Equilibrium is never 350°C, and a time of contact of 1.25 seconds, and an important of the contact of 1.25 seconds, and an important of the contact of the conta

copper in the form of a wire net at 480°C, hydrocyonic and in the principal product if the was polymerised. Using oxcess of ammonia, the acetylone could be fairly completely con verted into hydrocyenio acid According to Bredig, Elod and Demme, the reaction is C2Ha+2NHa=2HCN+3Ha. The use of an

hydrocyanic acid, under conditions studied by Bredig, Elod and Demmo (Ic.). They found

Contact time	% ammonta	% of disappearing NH	
seconds,	unchanged.	83 HCN	83 N ₁
65	90	78 5 77 2	21 5 22 8
1·2 0 67	35 9 72 3	69 2 57-0	30-8 43-0

ethylene products of the CH, +NH,

as ammonia. A large part of the ethylene yielded methane, carbon and hydrogen.

(c) Methane and Ammonia give very poor yields of hydrocyanic acid at temperatures up to 1,100°C. in the presence of alumina as a catalyst: this catalyst accelerates instead the decomposition of the methane and ammonia, as is shown by the following results (Bredig, Elöd and Demme, Z. Elektrochem. 1930, 36, 991) with a mixture of equal volumes of the two gases:

Contact time seconds.	Temp.	%,NH ₃ decomposed.	% NH ₃ as HCN	% CH ₄ decomposed.
8·8	800	56·5	trace	0·0
8·0	900	88·2	1.8	15·1
7·4	1000	98·5	9.0	47·6
6·8	1100	100	6.0	85·6

On the other hand, Bredig and Elöd claim better results in G.P. 548798, 1927, when using ethane instead of methane on a mixed aluminasilica catalyst at 850°C., a 35% yield of hydrocyanic acid being obtained. Probably acetylene is first formed.

Poindexter (U.S.P. 1584137, 1926) describes the production of hydrocyanic acid from ammonia and methane-containing gas (e.g. natural gas) at reaction temperatures between 950° and 1,450°C., depending on the materials of construction and the form of the apparatus. Preheating of the reactants is advisable in order that the contact time at reaction temperature may be kept as short as possible. 45-61% conversions of ammonia into hydrocyanic acid were obtained when 2 volumes of natural gas and I volume of ammonia were passed in 1 second or less through a glazed fireclay tube at 1,200°C. or higher. The loss of ammonia by decomposition was 27-39%. Various methods of supplying the necessary heat to support the reaction are described, such as the use of a chequer work stove preheated by combustion of fuel gas, or of a bed of coke which is blown red-hot by an air blast and then used as the reaction medium, or the addition to the gases of an insufficient quantity of oxygen for the com-bustion of their carbon, followed by explosion of the mixture. Further details are given in U.S.P. 1387170, 1920, and 1562914, 1922, according to the latter of which the product gas contains 11% HCN and considerable quantities of finely divided carbon.

The non-catalytic reaction between methane and ammonia is recommended by Wheeler and others (B.P. 335585, 335947, 1929), who find that the reaction

CH₄+NH₃=HCN+3H₂

proceeds at high velocities without the aid of solid surfaces if the temperature is of the order of 1,400°C., whereas the unwanted decompositions of methane and ammonia are catalysed by the walls of the chamber. They obtain good HCN efficiencies by choosing conditions which do not favour these decomconditions which do not favour these decom-positions, recommending, for example, the use of 1,100°C., and U.S.P. 2000134, 1935, describes

acid based on the nitric oxide, 2.4% appearing | chambers with the smallest surface and the dilution of the gases with hydrogen and moisture (B.P. 353407, 1930). They also state that the decomposition of ammonia and methane is minimised by building the reaction chambers with refractories free from iron, and in such a way that the internal surfaces are smooth. In B.P. 349958, 1930, they describe a form of apparatus designed to facilitate the transfer of the heat required to maintain the reaction, consisting of reaction chambers of long, deep and narrow form placed between long and deep heating flues.

One further process may be mentioned here (Ruhrchemie A.-G., G.P. 630767) in which mixtures of ammonia and methane are heated to high temperatures (1,400-1,500°C.) in vessels of non-porous alumina refractory at reduced pressure and very low times of contact. These reaction conditions have previously been found to be favourable for the production of acetylene. from methane, and the patent claims that, in the presence of ammonia, they lead also to the production of outstanding yields of hydrocyanic acid. A gas mixture containing 2 volumes of methane and 1 of ammonia gave, with a contact time of 0.05 second at a temperature of 1,500°C. and a pressure of 45 mm., an exit gas containing 12% HCN in amount corresponding to a 65% yield on ammonia and 32% on methane, with simultaneous production of acetylene. These results are in harmony with the view of H. Küster (Brennstoff-Chem. 1931, 12, 329) that the first step in the synthesis of hydrocyanic acid from methane is the splitting up of CH4 molocules into CH: groups.

The greatest technical difficulty in the methane-ammonia process is the rapid supply of heat at very high temperatures through refractory walls to the reacting gases, and various methods of overcoming it have been suggested. Peters and Küster (Brennstoff-Chem. 1931, 12, 122) heated the gases at 20-40 mm. pressure by means of an electric discharge, and obtained good yields. With a mixture of ammonia and methane in the volume ratio of 1:7, for example, they report complete cyanisa. tion of the ammonia, the excess methane being largely converted into acetylene. Wheeler also describes the production of hydrocyanic acid by passing induction sparks through a mixture of methane and ammonia (B.P. 325860, 1928), the power consumption being 0.05 kwh. per g. of HCN, or 0.01 kwh. per g. of HCN when the gas mixture is preheated to 1,000°C.

Another method of supplying internally a large part of the heat required is to add oxygen to the reacting gases, converting the endothermic synthesis

$$NH_3+1\frac{1}{2}O_2+CH_4$$

= $+100+114.9 \text{ kg.-cal.}$

Such a process is described in B.P. 361004, 1931, and 451609, 1935, the combustible mixture being passed at a high speed through a platinum

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the passage of the preheated muxture through | Dewar (Froe. Roy. Soc. 1010, 100, 100) | Dewar (Froe. Roy. Soc. 1010, 100) | D discusses the mechanism of the catalytic exidation at a platinum gauze, and postulates the intermediate production of "nitrovyl" groups HNO by the oxidation of ammonia. In the presence of excess of oxygen (as in technical ammonia oxidation for nitric acid), " nitroxyl " is further oxidised to nitrie exide and water. but when methane is present the following reactions are said to occur:

If methane is used in large excess, most of the oxygen is used in producing earbon monovide and hydrogen, and the yield of hydrocyanic acid The reactions by which "nitroxyl" and then hydrocyanic acid are formed are stated to occur very rapidly at the surface of the catalyst At 1,000°C, using a methane ammonia air mixture in the proportions required for the above oxidation equation, 58-63% of the ammonia was converted into hydrocyanic acid, 8-12% into nitrogen, and 26-30% was undecomposed.

The reaction between nitric oxide and methane in the presence of alumina at high temperatures wes mentioned by Elod and Nedelmann (2. Elektrochem 1927, 33, 217), and B P. 446277 (Du Pont de Nemours and Co) protects the use of one or more metals of the platinum group as catalysts for the reaction between hydrocarbons and nitric oxide, the mixture being diluted with nitrogen, carbon dioxide, steam or oxygen

7. From Nitrogen and Hydrocarbons —In 1868 Bertbelot (Compt rend 1868, 67, 1141) discovered that acetylene and nitrogen combine under the influence of strong induction eparks to give hydrocyanic soid by the reaction

which is reversible. He had previously shown (Ann. chim. phys. 1863, [111], 67, 52) that scety lene is formed in the electric are between carbon electrodes in an atmosphere of bydrogen, and that hydrocarbons in general yield acetylene under the influence of induction sparks. Acetylene is, in fact, one of the equilibrium products in the gas surrounding a carbon are burning in hydrogen or hydrocarbon vapour, being present to the extent of 7-8% by volume, along with 1-25% methane, 0.75% ethane, and no other bydrocarbons (Bone and Jerdan, J.C.S. 1901, 79, 1042). Berthelot drew the general conclusion that hydrocyanic acid must be formed in the carbon are burning in mixtures of hydrogen and nitrogen, and when sparks are passed through mixtures of nitrogen and hydrocarbons A mixture of 10 vols of acetylene, 14-5 vols, of nitrogen, and 75 5 vols of bydrogen is suitable for the synthesis of hydrogyanic acid, since mixtures richer in acetylene deposit carbon The cyanogen compound observed when the are burns between carbon electrodes in an atmosphere of nitrogen is by acceptance or hydrogen being derived from the electrodes or ephere of nitrogen is bydrocyanic acid, the

of hydrogen and nitrogen is passed through a carbon tube heated externally by an electric arc. Hitten and Smith (B.P. 23835 of 1906; Trans. Amer. Electrochem. Soc. 1908, 13, 333) have shown that the temperature of the arc is not necessary, hydrocyanic acid being formed when a carbon rod is beated at 1,700°C, in a mixture of bydrogen and nitrogen H. v. Wartenberg (Z. anorg Chem. 1907, 52, 299) carried out quantitative measurements of the equilibrium and obtained the following results with a mixture of equal volumes of hydrogen and nitrogen:

Temp. °C. 1635 1752 1875 % HCN in gas at equilibrium 195 3 10 4 70

Walks (Annalen, 1906, 345, 353) obtained a maximum HCN concentration of 35% with the same gas mixture in an arc baving an estimated temperature of about 3,250°C. From scetylene mixed with twice its volume of nitrogen, Hoyermann (Chem.-Ztg. 1902, 28, 70) obtained s 60-70% conversion into hydrocyenic ecid on passing the mixture into the arc through tubular

carbon electrodes.

Mixtures of acetylene end nitrogen under 3 atmospheres pressure were exploded in a closed bomh by Gerner and Matsuno (J.C.S. 1921, 119, 1903), and the cooled gases contained 1.57% HCN with an initial mixture containing 3% No. and 3 24% HCN with a mixture containing 22% No. The effect of increasing the temperature of the explosion by edding oxygen to the mixture was investigated by Garner and Saunders (J.CS. 1924, 125, 1634) The tem peratures attained in the explosion varied from 2,950°C. to 4,287°C, but the quantity of bydroevanue acid formed in the products corresponded with an equilibrium frozen at some lower temperature. Writing the reaction as

H_+2C+ N_=2HCN-55 4 kg.-cals.,

and using the relation

log kp= 13.210

+0 625 log T ~0 0009516T+ 1·18T³+34,

where $L_p = [HCN]^2/[H_s][N_s]$, for the variation of the equilibrium with temperature, it was calculated that the temperatures at which equilibrium was frozen lay between 1,940° and 2,282°C.

A considerable amount of work has been done on the synthesis of hydrocyanic acid from mtrogen and methane (or its equivalent) Gruszkiewicz (Z. Elektrochem. 1903, 9, 83) found that carbon menoxide, bydrogen and natrogen, maxed in the volume ratio of 2:1:1, gave amall amounts of hydrocyanic acid when exposed to induction sparks between platinum points. Lipinski (Z. Elektrochem. 1911, 17, 761) passed mixtures of methene, nitrogen, and hydrogen through a 2,200 volt alternating current are between platinum electrodes No carbon was deposited from a mixture containing 20% CH, 10% H, and 70% N, and under the

kwh. According to H. Philipp (Chem. and Met. Eng. 1920, 22, 313), Lipinski's process was tried on a commercial scale in 1914 at Neuhausen, and a yield of 30 g. HCN per kwh. attained. Briner and Baerfuss (Helv. Chim. Acta, 1919, 2, 663) report a yield of 7.39 g. HCN per kwli. on circulating a mixture of 5 parts of nitrogen and I of methane through an are at 505 volts between platinum electrodes 5 mm. apart.

A commercial process for the arc synthesis of hydrocyanic acid from methane and nitrogen has been developed by Soc. Chem. Ind. in Basle and H. Andriessens (B.P. 296355, 1927). Methane or petroleum vapour and nitrogen are circulated through an arc flame, producing a mixture of hydrocyanic acid and acetylene. The hydrocyanic acid is dissolved in caustic soda and worked up for sodium eyanide, and a large part of the acetylene is removed by solution in water under pressure before returning the gases to the arc (B.P. 195239). Küster (Brennstoff-Chem. 1931, 12, 329) carried out arc experiments under reduced pressure, and at the highest energy input was able to convert 80.5% of the methane into hydrocyanic acid and 13.9% into acetylene, starting with a mixture eon-taining 86.5% N₂ and 13% CH₄. With less nitrogen in the starting gas, the HCN/C₂H₂ ratio in the product fell. It is suggested that the methano is first decomposed into hydrogen and CH: groups, which react with activated nitrogen. A large oxcess of nitrogen and a high energy input are necessary to make the concentration of active nitrogen sufficient to react with all the CH: groups produced.
I.G. Farbenind. A.-G. (B.P. 294494, 1927) has

devised various modifications of arcs of the Schönherr type to make them applicable to mixtures of hydrogen, nitrogen and hydrocarbons, obtaining a lengthening of the arc by interposing between the central electrode and the earthed tubular counter-electrode a mantle of insulating material which surrounds the arc and lengthens its path. From mixtures containing 7-30% CH₄, yields of 50-65 litres of acetylene and 15-18 g. HCN per kwh. are

obtainable.

Properties of Hydrocyanic Acid.—Hydroeyanic acid does not occur in the free state in nature, but its compounds are widely distributed in plants. These compounds are glycosides which decompose, under the hydrolytic action of enzymes in the living cells, into hydrocyanic acid, a ketono or an aldehyde (e.g. acetone, benzaldohyde, ete.) and a sugar. For instance, the glucosido amygdalin, which is contained in bitter almonds, hydrolyses under the influence of the enzyme emulsin to give hydrocyanie acid, benzaldehydo and d-glucose; the hydrolysis is also offected by boiling with dilute acids. In rare cases a glycoside is found without an accompanying enzyme, as in the leaves of the common elder (Sambucus nigra) which contain the glucosido sambunigrin, isomeric with amygdalin (v. Cyanophoric glycosides). The quantity of hydrocyanic acid present varies in different parts of the same plant, and depends on the age of the plant, usually being greatest in young plants. It Voerkelius found that the NH3/HCN ratio

best conditions the HCN yield was I.75 g. per affected by climate and rainfall, and has been found to be increased by the use of nitrogenous fertilisers. The quantity present rarely exceeds 0.2% of the weight of the plant, and typical figures are given below:

> Great millet (Sorghum vulgare), 0.013-0.044% in young plants; an exceptional North American specimen yielding 0.114%.

Wild eherry (Prunus serotina), 0.09%. Cherry laurel (Prunus Laurocerasus), up to

0.2% in young leaves. Cassava, sweet (Manihot palmata), 0.016% in leaves, 0.043% in peel of stem, 0.005% in edible

part of root.

Cassava, bitter (M. utilissima), 0.041% in adible leaves, 0.113% in peel of stem, 0.053% in edible

part of root.

Rangoon beans (Phaseolus lunatus), 0.025% Linseed (Linum usitatissimum), 0.02-0.038%. Nandina domestica, 0.07-0.147%.

(Prunus communis, Bitter almonds amara), 0.12-0.18%.

Cases of eattle poisoning are not infrequent in hot countries where sorghum is used as fodder, and linseed meal and white clover are also sometimes poisonous. By thorough boiling with water the enzymes are destroyed and the cyanogenotic glucosides rendered harmless.

Minute traces of hydrocyanic acid have been observed in the products of incomplete combustion, such as the luminous flame of the Bunsen burner, boiler flue gases and tobacco smoke. Hydrocyanic acid is also always present in the products of the high-tomperature dry distillation of nitrogenous organic substances, and in this case is probably chiefly formed by tho action of ammonia on carbon:

$NH_3+C=HCN+H_2$

This reaction was used by Clouet (Ann. Chim. 1791, 11, 30) in the first synthesis of hydrocyanio Bergmann (J. für Gasbeleuchtung, 1896, 39, 117), Lance (Compt. rend. 1897, 124, 819), Voerkelius (Dissertation, Hannover, 1909), Carpenter and Linder (Alkal: Works, 45th Carpenter and Emide (211112), Report, 1908, 26; 46th Report, 1909, 21), and Winter (Bronnstoff-Chem. 1931, 12, 329). The Küster (Brennstoff-Chem. 1931, 12, 329). formation of hydrocyanic acid begins at 700°C.. and the best yields (calculated on ammonia) are obtained between 1,000° and 1,100°C. Tho reaction is reversible, but equilibrium conditions are never reached owing to the simultaneous decompositions of ammonia hydrocyanic acid at the temperatures in question. The composition of the product in any given experiment depends, therefore, on the relative rates of the main carbon-ammonia reaction and of the two decomposition reactions under the experimental conditions. Porcelain and iron, for example, favour HCNdecomposition, but little decomposition occurs with charcoal or when a layer of carbon has been deposited on the reaction tube. The rate of decomposition of the ammonia varies with the type of charcoal, being greater with more finely divided and with fresh samples of wood charcoal. is also dependent on the conditions of growth as I in the product was not greatly influenced by the

reaction time, but that ammonie decomposition action of formeldehyde and hydroxylamine, en increases with the reaction time. Küster experi-heating the two compounds alone or with increases with the reaction time. August a phosphorus pentoxide (Dunstan end Bossi, coke oven a mixture of nitrogen and hydrogen J.C.S. 1898, 73, 353), and (b) the combination coke oven, a mixture of nitrogen and hydrogen in the volume retio of 1:3, and containing 0 2% of ammonia, being passed over coke. The results were:

Temp C. % NH, giving HCN 600 700 B00 900 1000 1100 25 1 40 0 128 % NH, decom 53 6 85 42 B 63 3

Nernst'a heat theorem has been applied to this reaction by you Wartenberg (Z. anorg. Chem. 1907, 52, 299) with the following results:

800 1400 Temp. °C. 500 1100 0 27 0 003 0 0002

[PNH,/(PH,×PHCN)] 632 The equilibrium quantity of hydrocyanic acid increases merkedly with rise of temperature, the heat of the endothermic reaction being -39,500 cal. per mol. of HCN. These figures indicate that equilibrium was never even approximately attained in Voerkehns' experiments, and the formation of appreciable quantitles of hydrocyanic acid at 600°C, in Küster's experiments suggests that reactions between

ammonia and hydrocarbons may contribute.

Hydrocyanic acid was formerly prepared by
distilling potassium ferrocyanide with dilute sustaining poeassium retrocyanide with dilute sulphurio acid (see under Technical Methods of Preparation, p. 492), hat is nowadays more conveniently prepared from codum cyanide solution and relatively concentrated authoric acid, a mathod first described by Wade and Panting (J.C.S. 1898, 73, 255). A 30-35% A 30-35% solution of sodium eyenide is run into hot 759 aulphurio acid, using 11 mols. of sulphuric acid to 2 mols. of cyanide The cyanide solution should he allowed to enter the acid just below ita surface, in order to avoid contact of the alkaline cyanide solution with hydrocyanio acid vapour, which encoutages polymerisation of the HCN. The vapour may be dried by means of calcum chloride at 30°C, and then condanged in en ice cooled vessel. It may be purified from tracee of water and ammonia by distillation over phosphorus pentovide (Nef, Annalen, 1895, 287, 265) Details of a similar laboratory procedure, adding a small amount of ferrous sulphate to the reaction mixture, are given hy Slotta (Ber. 1934, 67, [B], 1028)

The above process is frequently used for the menufacture of hydrocyanic ecid (Pelton and Schwarz, Chem. end Met. Eng. 1919, 20, 165; Carlisle, Ind. Eng Chem. 1933, 25, 959). The evanide sulphuric acid reection mixture is distilled to give an aqueous distillate of 80% atrength, which is acidified to inhibit decomposition, and rectified by redistillation. B.P. 401351, 1931, proposes the use of eulphur cloxide as a gaseons etabliser to prevent decomposition throughout the reaction and of this antidote is due to the conversion of the distillation systems; it may conveniently be hydrocyenic acid into ferrocyenide, which is generated by edding sodium sulphite to the relatively non toxic. A summarised description cyenido befors the reaction with the acid.

Other methods of preparing hydrocyanic various antidotes for hydrocyanic edit possible with a restrict of the property of the an isomer of formamide obtained by the inter

of hydrogen and cyanogen by heating the mixture to 500-550°C. (Berthelot, Ann. chim. phys. 1879, [5], 18, 380).

Hydrocyanic coid is a colourless liquid having a characteristic emell, or more correctly taste, which produces a choking constition in tha larynx. It is one of the most poisonous sub-stances known; the fatal dose for e human being is approximately 1 mg, per kg, of body weight, or 006-007 g. for a full grown man. It may be introduced into the body hy inhalation of the vapour or by absorption of the hauid through the ekin, and its lethal action is very rapid, large doses causing almost instantaneous death. On the other head, air containing minute concentrations of the vepour, but in which its emell is distinctly perceptible, may be hreathed constantly without ill effects. Expen ments with cets, for instance, showed that con centrations of 0 003-0 001% could be tolerated, hut that 0 005% produced symptoms of poisoning. It appears that the metabolism is shie to convert hydrocyanic ecid into relatively non toxic thiooyenete at a limited rate, and if the rate of supply of hydrocysnic acid is lower than this rate, no lethel effect is produced cyanate has been detected in the urine of dogs which had been treated with regular small doses

of potassium cyanide. The first symptoms of poisoning are giddiness and darkening of the visuel field; these symp toms disappear after a few minutes' exposure to fresh air. They are followed by unconsciousness accompanied by muscular cramps and more or less complete cessation of respiration So long as the heart is still acting, however, artificial respiration and the use of oxygen (or a mixture of oxygen and carhon dioxida) may be successful at this stage in inducing recovery. Heert stimulants such as anyl nitrite (applied by inhelation of the vapour), lobeline, and coramine (applied by intravenous injection) may be applied by e doctor in desperate cases. Other antidotes which have recently been the subject of reasarch are various sulphus producing preparations, such as colloidal sulphur, thiosulphate, and tetrathionate, sodium nitrite and mixtures of nitrite with thiosulphate,

methylene blne, and oxidising egente such as hydrogen peroxide and permanganate When a cyanide has been swallowed, hydrocyanic acid is liberated in the atomach by the acids in the gastric juicee, and in this cass the simplest and most effective antidote is prepared by mixing colutions of ferrous sulphate and sodium carbonate. It is important that the sodium carbonate should he in excess so that the mixture is alkaline; the detoxifying action of the mode of ection and relative efficiency of

The poisonous ection of hydrocysnic acid

appears to consist partly of a direct paralysing effect on the respiratory nerve centres and partly of an inhibitory action on that enzyme of the red-blood corpuscles which brings about the transference of oxygen from oxyhæmoglobin to the oxidisable substances in the body: the venous blood therefore becomes arterial, and death is due partly to a kind of internal suffocation. The suggestion that the poisonous action of hydrocyanic acid might be due to its inhibitory action on the oxidising power of the blood was first made by Schönbein (J. pr. Chem. 1868, [i], 105, 202), who showed that the acid inhibits the catalytic decomposition of hydrogen peroxide by plant ferments, and also by the redcorpuscles. Later Bredig and Berneck (Z. physikal. Chem. 1899, 31, 329) showed that it has the same effect on the inorganic catalyst, colloidal platinum. The ferments recover their activity when the hydrocyanic acid is removed. This property has been applied in the preservation of fruit.

The physical properties of liquid hydrocyanic

acid are tabulated below:

Density (Fredenhagen and Dahmlos, Z. anorg. Chem. 1929, 179, 77):

Temp. °C. −13 Density 0.7326 0.7150 0.7096 0.6950 0.6884

Freezing-point (Tammann, Ann. Physik. 1899, 68, 576):

Pressure (atm.) 500 4,000 Freezing-point (°C.) -13.4 -2.88+50.1

Vapour Pressure (Bredig and Teichmann, Z. Elektrochem. 1925, 31, 449):

Temp. (°C.) -15.2-1010.8 Vap. press. (mm.) . 110 165 256427 Temp. (°C.) 14.8 18.0 25.6 29.8 Vap. press. (mm.) 504 567 760 888 Temp. (°C.) 100 64.8 78.5 39.43.71 5.35Vap. press. (atm.) 1.63

Thermal Data.—Heat of formation of liquid from its elements (amorphous carbon) -21.8cal. per mol. (Thomsen); (diamond) -24.8 cal. per mol. (Berthelot). Latent heat of vaporisation at 25°C., 246.8 cals. per g. (Perry and Porter, J. Amer. Chem. Soc. 1926, 48, 299)

Heat of solution of liquid acid in water: according to Busay and Buignet, when 1 g.-mol. of liquid HCN is mixed with 11 g.-mols. of water, a fall of temperature of 9.75°C. occurs.

Aqueous Solutions.—The specific gravities of aqueous solutions have been measured by Walker and Marvin (Ind. Eng. Chem. 1926, 18, 139), whose results are given in an abbreviated form below:

Sp.gr.	0°	5°	10°	15°	20°	25°
0.682	_					100
0.690					99.8	97.7
0.700	_			99.0	96.9	94.8
0.710	_		98.1	96.0	93.9	92.0
0.720	99.3	97.2	95.1	93.0	91.0	$89 \cdot 1$
0.730	96.2	94.2	92.0	90.0	88.0	86.2
0.740	93.1	91.1	89.0	87.0	85.1	83.3

These data are often used to estimate the percentage of water in the commercial acid.

The eutectic point, ice+solid HCN+liquid phase containing HCN, is -23.4°C. 74.5 molar percentage of

The dielectric constant of liquid hydrocyanic acid is 95, the largest known with the exception of that of hydrogen peroxide (Schlundt). The acid is miscible in all proportions with water, alcohol, ether, and many other substances, but it is almost insoluble in light petroleum. It is itself an excellent solvent for many solid substances; among inorganic salts the iodides are usually readily soluble. The solutions are good conductors of electricity, the conductivity of solutions of potassium iodide being about 3½ times greater than that of the corresponding aqueous solutions.

Hydrocyanic acid is one of the weakest acids, its dissociation constant in aqueous solution at 18°C. being 1.3×10⁻⁹, or 44 times smaller than that of sulphuretted hydrogen and 230 times smaller than that of carbonic acid (Walker and Cormack, J.C.S. 1900, 77, 16). In a 0·1 N solution, only 0.011% of the acid is ionically dissociated, so that it does not behave as an acid towards indicators, and its salts are very largely hydrolysed in aqueous solution. The hydrolysis of alkali cyanides has been studied by Harman and Worley (Trans. Faraday Soc. 1925, 20, 502), who measured the dissociation constant k = [KOH][HCN]/[KCN] of the KCN+H₂O ⇌ KOH+HCN reaction various temperatures with the following results:

Temp. °C. $k \times 10^4$. 0.179 0.2290.2540.304

The same figures apply to sodium cyanide. In the concentration of an aqueous cyanide solution free hydrocyanic acid produced by hydrolysis escapes until a sufficient excess of the base has accumulated to stop the hydrolysis: with stronger bases, such as caustic soda, the loss of hydrocyanic acid is slight, but with weaker bases it is considerable. In the case of volatile bases (e.g. ammonia) and insoluble bases (e.g. calcium hydroxide), the continued removal of the base from the solution (by volatilisation or precipitation) prevents the attainment of equilibrium, and nearly the whole of the hydrocyanic acid escapes. The carbon dioxide of the atmosphere will displace hydrocyanic acid from aqueous cyanide solutions, so that such solutions in confined spaces may develop dangerous HCN concentrations in the surrounding air.

Pure hydrocyanic acid, which has been dried over calcium chloride or distilled over phosphorus pentoxide, can be kept unchanged for long periods at the ordinary temperature. Instability is induced by heating the liquid to 100°C., or at the ordinary temperature by the presence of a small amount of water or of alkaline impurities such as ammonia or sodium cyanide, and the liquid gradually darkens in colour through yellow and brown, being finally converted into a non-volatile black solid of about the same empirical composition as HCN (Walker and Eldred, Ind. Eng. Chem. 1925, 17, 1074). This black polymer is itself a catalyst for the polymerisation. The decomposition occasionally proceeds with explosive violence owing to the

rapid rise of temperature which accompanies the | England, hydrocyanic acid is being used for the reaction, and Salomone (Gazzetta, 1912, 42, (i), eradication of vermin. 617) isolated from the products of an explosion polymer can be separated by extraction with aminomalome acid, NH₂CH (CN)₂, since it 25, 959; U.S.P. 1352685, 1355384, 144430, gives carbon dioxide, ammonia, and aminoacetic acid on heating with baryta water. From the same material, Bedel (Compt. rend. 1923, 176, 168) obtained a tetramer which he helieves to be the hydrocyanide of amino-methane dinitrile. The aqueous and also becomes brown in the presence of alkah eyamdes and deposits a dark brown amorphous anbitance, azulmin or azulmic acid, containing carbon, nxygen, nitrogen, and hydrogen in variable proportions. Whenever water is present, carbon monoxide and ammonia are found among the decomposition products. The discoloration of solid alkalı cyanide on exposure to air is due to the hydrocyanic acid liberated at its surface undergoing the change to azulmie acid

The commercial acid, which generally contains about 2% of water, is stabilised to inhibit decomposition by the addition of small quantities of strong acids, such as sulphure acid or phosphoric acid, or better, oxsho acid, which is less corrosive to iron containers (G.P. 352979, 1919). Organic compounds which are readily hydrolysed to acid products are also suitable, eg. cyanogen chloride (BP 258324, 1925),

chloroformic esters (B P. 254747, 1925), The only useful application of hydrocyanic acid la in pest control, where its poisonous action is used against insects on growing plants and in various stored products, and against rodents and other pests in the furnigation of ships and mills, warehouses and other buildings, and in quarantine operations. This application was first developed by the Agricultural Experiment Station of the University of California for the fumigation of citrus trees (D. W. Coquillet and H. D. Bishop, California Station Bull. 122). The acid vapour is applied by erecting a tent over the tree and generating the acid maide it. The dosage applied must be high enough to kill the scale insects within about half an hour, but not high enough to cause damage to the foliage. Allowance must also be made for loss of fnmigant by diffusion of the vapour through tha tent during the period of exposure. Foliage is less hable to injury at night than in sunlight, and fumigation should not be carried out in damp conditions. The safe concentration of hydrocyanic acid depends on the kind of tree and on cyanic acid depends on the annu or irre and on the time of the year. The minimum lethal dependence waries widely for different species of pests, green aphis requiring only 2 g. of HCM per 1,000 cu. ft. of air space, and red appier up to 11 as also used for the synthesis of aromatic 30 g per 1,000 cu. ft. in greenhouse fumigation. aldehydea by Gattermann's reaction. The use of hydrocyanic acid in the fumigation of ships, warehouses and mills is now well The constitution of hydrocyanic acid is a very established in many parts of the world, and is complex question, and cannot be said to have also extending to the treatment of stirred been definitely settled. The behaviour of its products. In alum clearance softeness in metallice saits would lead to contradictory

In early fumigation practice the hydrocyanic colourless crystals of a substance having tha acid was generated in situ by the addition of composition of cyanic acid, the molecular weight solid acduum cyanide to a mixture of aulphune of which in henzene solution corresponded to acid and water, but since 1917 houd hydro-(HCNO), From the solid black HCN cyanic acid of about 98% strength has been polymer mentioned above, a termolecular marketed in tinned iron drums or steel cylinders (see Pelton and Schwarz, Chem. and Met. Eng. ether (Nef), which is believed to he tha nitrile of 1919, 20, 165; Carlisle, Ind. Eng Chem 1933.

tion the acid may be absorbed in granules of kieselgubr previously freed from alkaline impurities (B.P. 271236, 1926), or in discs of wood pnlp. As already mentioned under Calcium Cyanide, p. 492, powdered calcium eyanida may also be used in a moist atmosphere for the generation of hydroeyanic acid. Hydrocyanic acid readily forms addition

products; for instance, with hydrochloric and in ethyl acetate solution it gives the compound 2HCN-3HCl, which has been shown to be dichloromethylformamidine hydrochloride,

HN CH NH CHCI2, HCI.

This compound decomposes at 100°C, into hydrochlorie acid and a volatile substance 2HCN-HCl, which is chloromethylene forms-midine, HN-CH-N.CHCl. The latter compound is also formed when dichloromethylformamidine hydrochleride is treated with excess of liquid hydrocyanic acid at 40°C. When warmed with quincline, chloromethylene formamidine gives a quantitative yield of iminoformyl sectyanide HN CH NC, a dimer of hydrocyanic acid (Hinkel and Dunn, J.C.S. 1930, 1834), which is a white solid melting et 85°C. Hinkel, Avhor and Bevnon (1814) 85°C. Hinkel, Ayling and Beynon (itid 1935, 674) have investigated the hydrolysis of this compound with cold water to give ammonium formate, and its formation of insoluble complex compounds with silver mitrate, mercurio chloride and aluminum chloride. Hydrocyanio acid itself forms a compound with aluminium chloride, having the composition AICI, 2HCN (Hinkel and Dunn, total. 1931, 3343), which is stated to be identical with the aluminium chloride com-pound of the dimer. The compound is unstable in moist air, and slowly evolves hydrocyanic acid in racus or on heating to 100°C

Hydrocyanic acid is often used for the synthesis of a hydroxy acids from aldehydes and ketones, with which it combines under the infinence of alkaline catalysts to give cyanhydrins, which are the nitriles of a hydroxyacids, e g.:

CH₂CHO+HCN → CH₃·CH(OH)(CN) Acctaldehyde

CONSTITUTION OF HYDROCYANIC

esters to give nitriles R.C.N as principal product, whereas silver cyanide yields carhylamines or isocyanides R.N.C. The force of this evidence is in any case considerably reduced by the fact that in neither case is the one product exclusively formed. Other properties point to a difference in structure hetween the alkali cyanides and those of the heavy metals; for instance, silver and mercuric cyanides cannot be oxidised by permanganate, whereas potassium cyanide gives cyanate. On the other hand, it should he remembered that the heavy metal cyanides are scarcely ionised at all and their reactions are therefore those of the undissociated molecules; the alkali cyanides, on the other hand, are very largely ionised, and their reactions are those of the ions. For this reason it may be concluded that all metallic cyanides have the same constitution, and the generally accepted one is the isonitrile configuration M·N:C. This agrees with their ready formation of stable addition compounds, in which they resemble the isonitriles and differ from the nitriles.

Hydrocyanic acid itself may he tautomeric, $H \cdot N : C \rightleftharpoons H \cdot C : N$, but the weight of evidence in this case supports the nitrile formula as being at any rate greatly predominating. Like the nitriles, hydrocyanic acid is hardly affected hy dilute mineral acids, from which it may he distilled unchanged; the carbylamines or isonitriles, on the other hand, are violently decomposed. With alkalis, hydrocyanic acid is readily hydrolysed or decomposed, but even boiling alkalis are without action on isocyanides Its high dielectric constant and marked solvent and dissociating powers relate it to the nitriles rather than to the carbylamines (Wade, J.C.S. 1902, 81, 1613). Wade is also of the opinion that its physiological properties lend support to its configuration as a nitrile, but Nef draws the opposite conclusion. Anhydrous hydrocyanic acid and the nitriles do not dissolve silver cyanide, whereas the isocyanides dissolve it

readily.

VI. CYANOGEN AND ITS DERIVA-TIVES.

Cyanogen, (CN)₂, is prepared by heating dry mercuric cyanide at a dull red heat, the reaction by which Gay-Lussac discovered it in 1815:

$$Hg(CN)_2=Hg+(CN)_2$$

It may also he prepared by the interaction of sodium cyanide and copper sulphate, when the first product is a complex cupricyanide, NaCu(CN)3, which decomposes on heating to give cuprous cyanide, $Cu_2(\hat{C}N)_2$, and cyanogen. A further yield of cyanogen may he obtained by treating the cuprous cyanide precipitate with ferric chloride solution of sp.gr. 1.26:

$$Cu_2(CN)_2+2FeCl_3$$

= $(CN)_2+Cu_2Cl_2+2FeCl_2$
By all these methods a part of the averagen is

By all these methods a part of the cyanogen is paracyanogen. Paracyanogen appears to be acid, and contains also oxalic acid and urea. the stable form at ordinary temperatures, but at Cyanogen dissolves in caustic alkalis to give

conclusions: potassium cyanide reacts with | high temperatures is converted into cyanogen. The velocity of the change is very small at temperatures helow 300°C., so that cyanogen gas can he kept without change in the cold. The polymerisation of cyanogen, and the catalytic effect of glass, quartz, potassium cyanide, and sodamide have heen studied by Perret and Krawczynski (Bull. Soc. chim. 1932, [iv] 51, 622).

The direct synthesis of cyanogen from its elements is not of practical importance hut has theoretical interest in relation to several technical cyanide syntheses. Cyanogen is formed when the electric arc hurns between pure carhon electrodes in an atmosphere of nitrogen, though it cannot he detected in the gaseous product withdrawn from the arc (Wallis, Annalen, 1906, 345, 353). Nernst's heat theorem would indicate that considerable quantities of cyanogen exist in equilibrium with carhon and nitrogen at the temperature of the

Temp. (°C.) % (CN)₂ in gas at 2,000 2,500 3,000 3,250 equili brium 17.0 45.0 0.01

and von Wartenberg (Z. anorg. Chem. 1907, 52, 299) observed a very intense cyanogen spectrum in the arc gases, and found that carhon is deposited at some distance from the arc. Cyanogen is therefore essentially unstable at temperatures below 2,500°C., and can only be preserved at much lower temperatures on account of its slow rate of decomposition. It can be detonated hy means of mercury fulminate, and on heating its rate of decomposition hecomes measurable at 1,200°C. (Berthelot, "Force des Matières Explosives," Paris, 1883, p. 113). It may be concluded from these figures that the direct union of carbon and nitrogen to cyanogen can play no part in the formation of cyanides by processes operated at ordinary furnace temperatures. On the other hand, it should he mentioned that cyanogen is said to be produced when nitrogen is passed over iron covered with coke at 1,500-1,800°C., and Belg. P. 378694, 1931, protects such a process, in which nitrogen at 50 atmospheres is heated in the presence of reduced iron with compressed carbon monoxide. Presnmably there is inter-mediate formation of iron carbide and iron nitride.

Cyanogen is a very poisonous gas which hurns with a characteristic peach-blossom-coloured flame. The liquid has a density of 0.89 at 13.8°C., hoils at -20.7°C. at 1 atmosphere, and freezes at -34.4°C. Its vapour pressure at -22.9° is 701.6 mm., and at -14.4° is 1030.4 mm. (Cook and Robinson, J.C.S. 1935, 1001). The critical temperature and pressure are 128.3°C. and 59.6 atmospheres respectively (Cardoso and Baume, Compt. rend. 1910, 151, 141). Water dissolves 4.95 volumes of the gas at 0°C. At 18°C. and higher the dissolved cyanogen is hydrolysed to hydrocyanic acid and cyanic acid, the latter being further hydrolysed to give carhon dioxide and ammonia (Naumann, Z. Elektrochem. 1910, 16, 772). After some always polymerised to dark hrown, non-volatile time the solution deposits dark hrown azulmic

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cyanide and cyanate hy a reaction analogous the cold asturated solution containing 6% of to those of the halogens:

CNCI. The heat of formation of the house

Silver nifrate solution has no action on cyanogen which may be separated from a mixture with hydrocyanic acid by passing through salver nitrate solution to remove the HCN, and may be estimated by treatment with caustic sodi and titrating the cyanide formed (Walls, Annalen, 1906, 345, 353).

Cyanogen is held to have the nutrile structure N;C—C;N, and not the isomitrile structure C:N—N.C or a mixed atructure N;C—N.C, becausa of its formation by the dehydration of ammonium oxelate or oxemide with phosphorus pentoxide:

which shows that the two carbons are linked together.

Cyanogen chloride, CNCI, is prepared by the action of chlorine on various metallic cyanides. Heavy metal cyanides, such as msrcuno (Ssrullas, 1827) and zino (Held, 1897) may be used, but the use of aqueous hydro-cyanic acid, discovered by Berthollet in 1787, is now a preferred method. Alkali cyanides are not suitable because they react violently with cyanogen chlorids in any hut cold and very dilute solutions, giving dark hrown amorphous products. U.S.P. 1938324, 1933, however, describes the production of cyanogen chloride from chlorine and sodium cyanide in a mixture of carbon tetrachloride and glacial acetic acid in the absence of water. For the preparation from aqueous hydrocyanis acid (Price and Green, J.S.C.I. 1920, 39, 987), a 12-15% solution of pure hydrocyamo acid is treated with a stream of finely divided chlorine bubbles, obtained by injecting the gas through a porous earthenware plate at about 40°C. The cyanogen chloride distils off as it is formed together with 10-20% of hydrocyanic acid. It is purified by retreat-ment with chlorine, or by shaking with zine oxide to remova the hydrocyamo acid almost completely as zinc cyanide. Hydrochlorie acid in the product a removed by distilling the liquid over sodium hicarbonate. A technical process for the preparation from aqueous hydrocyanic acid and chlorina under pressure is described in B.P. 347989, 1929.

Cyanogen chloride is a colourless liquid of dansity 1-193 at 14°C. (Cook and Robinson, J.C.S. 1935, 1001), which sobdifies at -8°C. Its vapour pressure at various temperatures was measured by Ragnatt;

Its boiling-point at 760 mm. is 12 66°C. The vapour, which is very peisonous, attacks the ayes violently. It is alightly soluble in water,

the cold asturated solution containing 6%, at CNCI. The heat of formation of the loyal from its elements is -26 9 cal. per mol. It is stable for an indefinite period when pure, but in the presence of hydrochloric acid it polymeries to cyanuric chloride, C₂N_CI₁, Aqueon hydrochloric acid (stronger than 2N) hydrolyne it thus:

Its behaviour towards alkalis is the sams as that of cyanogen bromida (see below), and indicates that cyanogen chloride is the acid chloride of cyanic acid, CNOH. Hydriodic acid hardly attacks it in tha cold, and it is without actors on iron and copper in the cold.

Cyanogen bromlde, CNBr, may be pre-pared by the action of hromine on a metallic cyanide or on hydrocyanic acid. Slotta (Ber. 1934, 67, (B], 1028) gives details of the action of sodium cyanida solution on bromma in a well atured flask immersed in a freezing-mixture, the bromocyanogen being distilled direct from the reaction mixture through a layer of calcum chloride into a cooled receiver. A laboratory preparation is also described in "Organs Syntheses," 1931, 11, 30). The compound so technical importance because its addition to a cyanide solution causes more rapid dissolution of gold (Sulman and Teed, B.P. 18592 of 1894; J.S.C.I. 1897, 16, 981). It is used in the treatment of a few refractory ores, especially those in which the gold is associated with tellurides, as in Western Australia. For this purpose it is always manufactured at the place of use by the cotice of the place of use by the cotice of the place of the place of the cotice of the cot lace of use hy the action of acidified bromidebromate solution on a concentrated solution of sodium cyanide; other oxidising agents may be used instead of hromate. Since the action of hromine on sodium cyanide also produces sodium bromide, the addition of a further quantity of bromate will yield a further supply of bramus and hence of cyanogen bromide; with an adequate quantity of hromate, therefore, the reaction is (Göpner, Z. angew. Chem. 1901, 14, 355):

2NaBr+NaBrO₃+3NaCN+3H₂SO₄ ≈3Na₂SO₄+3CNBr+3H₂O

The reaction countring when gold dissolves in bromocyanide adution is as follows:

2Au+3NaCN+BrCN

=2NaAu(CN)₂+NaBr Ite effact is thus seen to be dus to its character as being both a cyanida compound and an

madaing agent (see Gold Cyandides, p. 480).

Cyanogan bromidio as colouries, crystallae
anbatance melting at 52°C, and bouling at 6°C.
Its solubility in water is about 5-6°% at the
ordinary temperatura. It is very poisonous, adhas an intensity irritating effect on the innecess
membrane and the eyes. Lifts cyanogathorids, the pure substance and the bydrobronic
acid cause its rapid polymerisation to cyanorie
bromide. C., N., Er, with simultaneous partial
decomposition. It is very alowly hydrolywed
water to hydrobromic and copan acides:

CNBr+H.O-HCNO+HBr,

occur instantaneously with caustic alkalis, much more slowly with alkali carbonates, and not at all with bicarbonates. In connection with the bromo-cyanide process of gold extraction, it is of interest to note that cyanogen bromide slowly reacts with potassium cyanide in aqueous solution, giving cyanogen:

$CNBr+KCN=(CN)_2+KBr$

A 0·1-0·2% CNBr solution is completely decomposed in 3-6 hours by KCN solution of less than 1% strength.

In aqueous solution, cyanogen bromide acts as an oxidising agent, converting hydriodic acid into iodine, and sulphuretted hydrogen into sulphur:

In these reactions the bromine behaves like hypobromous acid, which is typical of bromine attached to nitrogen (Chattaway and Wadmore, J.C.S. 1902, 81, 191), so that the compound must be formulated as an iso-cyanogen derivative, C:N·Br. The reaction with potassium iodide provides a convenient method for the quantitative estimation of oyanogen bromide.

Cyanamide, NH₂·CN, is obtained by the action of ammonia on an ethereal solution of cyanogen chloride (Cloëz and Cannizzaro, Compt. rend. 1851, 32, 62), when ammonium chloride crystallises out and cyanamide is recovered from the ethereal solution by evaporation. It is also obtained by the action of

mercurio oxide on thiourea:

$$CS(NH_2)_2+HgO=NH_2\cdot CN+HgS+H_2O$$

Several guanidine derivatives yield cyanamide on boiling with water or alkalis; for example, nitroso-guanidine decomposes thus (Thicle, Annalen, 1893, 273, 136):

Cyanamide is most easily prepared from its metallic salts. Disodium cyanamide, Na, NCN, is formed by the action of sodamide on carbon or on sodium cyanide (see Alkali Metal Cyanides, p. 480), or by passing carbon dioxide over heated sodamide (Geuther, dioxide Annalen, 1858, 108, 93):

NaNH₂+CO₂=NH₂·COONa

=NaCNO+H₂O

NaCNO+H,NNa=Na,N·CN+H,O

It is also formed when either sodium cyanate or sodium cyanide is fused with sodium hydroxide, or when ammonia reacts with sodium carbonate at 500-700°C. (G.P. 582625, 1928). The calcium and barium salts are produced by the nitrogenation of the carbides (see under Alkaline Earth Cyanides, p. 487). and by several other reactions at moderate temperatures where cyanide would be expected, e.g. from lime and hydrocyanic acid (Franck and Bank, Z. anorg. Chem. 1933, 215, 415) $= H_2N(C:NH)NH\cdot CN + H_2O$ $= H_2N(C:NH)NH\cdot CO$ and by heating the ferrocyanides. They are a strong base which neutralises the acid.

the cyanic acid being next hydrolysed to also formed by heating the carbamates or the ammonia and carbon dioxide. These reactions cyanates to redness (Drechsel, J. pr. Chem. 1877, [ii], 16, 188; 1880, [ii], 21, 77) (see also G.P. 591039, 1929):

> $Ca(OCONH_2)_2 = CaN \cdot CN + 2H_2O + CO_2$ Ca(OCN)₂=CaN·CN+CO₂

Cyanamide is easily prepared from the disodium salt by neutralising its cold, aqueous solution with nitric acid and evaporating to dryness in a vacuum; the dry residue is extracted with ether. According to Liebknecht (Angew. Chem. 1932, 45, 584; G.P. 601465, 1934), it is best to treat the disodium compound first with absolute alcohol to produce the monosodium salt and sodium ethoxide, and then to treat the mono-sodium salt with hydrogen chloride in ether solution. Werner (J.C.S 1916, 109, 1325) describes its preparation from the readily accessible calcium salt: 100 g. are mixed with 125 g. of acetic acid and 120 g. of water, the mixture well kneaded and left for 24 hours, after which the friable mass is powdered and extracted with other in a Soxhlet apparatus. The yield is 94-96% of the theorctical

Cyanamide forms colourless, deliquescent crystals melting at 43-44°C. (Colson, J.C.S. 1917, 111, 555). It is readily soluble in water, alcohol and ether, and sparingly soluble in carbon bisulphide, benzene and chloroform. On heating it polymerises very readily to dicyanodiamide, H₂N(C:NH)NH·CN, a liquid which boils at 140°C. under 19 mm. with decomposition to melamine. Pinck and Hetherington (Ind. Eng. Chem. 1935, 27, 834) produce the dimer in 98% yield from concentrated cyanamide solutions on heating with small amounts of ammonia. The neutral aqueous solution is stable, but polymerisation occurs in the presence of either acids or alkalis. The physico-chemical aspect of the process has been studied by Morrell and Burgen (J.C.S. 1914, 105, 576) and by Grube and Krüger (Z. physikal. Chem. 1914, 86, 65), who find that in alkaline solution the reaction takes place between undissociated cyanamide molecules and cyanamide ions:

 $H_2CN_2+HCN_2'=H_3C_2N_4'$

Hence the rate of polymerisation is at a maximum in a solution containing one-half of its cyanamide in the ionised form, which is virtually the case when one-half equivalent of a strong base is added to the aqueous solution. Weak bases, since their cyanamide salts are considerably hydrolysed, have a much smaller polymerising action than strong ones.

The action of acids on cyanamide causes polymerisation and hydrolysis (to urea) simultaneously. Cyanamide treated with 10-20% sulphuric acid at the ordinary temperatures is almost wholly converted into urea, whereas near the boiling-point polymerisation pre-dominates (Carlson, Z. angew. Chem. 1914, 27, 724). The reaction is further complicated by the hydrolysis of the dicyanodiamide to dicyanodiamidine,

=H2N(C:NH)NH·CO·NH2,

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Cyanamide readily forms addition com- conditions by reaction between carbon monovide pounds. For example, when dry hydrogen and ammonia (p. 494). An interesting synthesis chloride is passed into a dry ethereal solution, a for cyanic acid is described by Lauda (Compt. white crystalline aubstance baving the formula H, CN, 2HCl is formed (Mulder and Smit,

Ber. 1874, 7, 1634).

Cyanamide is an extremely weak acid, the first dissociation constant (H₂CN₂ ⇒ H⁺+HCN₂) being of the order of 10⁻¹¹ (Grube and Kruger, Z. physikal. Chem. 1914, 86, 65). The monoaodum salt, NaHCN₂, crystallises from a solu-tion of cyanamide in absolute alcohol when sodium ethoxide is added (Liebknecht, Angew. Chem. 1932, 45, 584); the disodium salt can only be prepared by one of the dry fusion methods mentioned above. The calcum salt, Ca(HCN₂)₂, can be obtained in solution by decomposing ordinary calcium cyanamide with water. The silver salt, Ag₂CN₂, is obtained as a canary yellow precipitate by adding silver nitrate solution to a dilute ammoniacal solution of cyanamide; it is soluble in cold, dilute nitrie acid. The lead salt is an orange yellow

precipitate,
Although there is some doubt as to the constitution of cyanamide (H, N.C: N or H, N N C), the balance of evidence is in favour of the normal cyanide structure, for cyans mide ahowa many reactions of a normal cyanide or nitrile, e g. reaction with water to give the acid amide ures, with sulphuretted hydrogen to give thio urea, and with ammonia to give the amidine guandine, HN:C(NH₂)₂. Its formation from cyanogen chloride, to which the sometrile atruc ture is assigned, can be explained by the intermediate formation of an additive compound

(Nef):

VII. CYANATES.

The cyanates, salts of the acid HCNO, are not themselves of technical importance, but derive interest from their close relationship to the cyanides, from which they are readily formed by oxidation, and into which they are converted by reduction. This hehaviour is held to support the " isocyanide " structure for the metallic cyanides, since the oxidation to cyanate is than represented as a transition from divalent to tetravalent carbon:

Na·N:C→Na·N C:O

In the case of erganic derivatives of cyame acid, two series of compounds should exist, the normal cyanates R O.C: N, and the isocyanates R N.C:O. Only the latter have been solated, however, as the normal cyanatea are so unstable soon as they are formed.

Several reactions in which cyanates are formed have already been described, and need only be solution in dry other is comparatively stable, briefly mentioned here. Cyanates are formed but if a small amount of triethyl phosphina is Ammonium cyanate is produced under certain (HCNO)z, where x prehably equals 3, has been

rend. 1932, 194, 2070), who oxidises carbon compounds with potassium permanganate in compounds wan possessian permanganate in concentrated ammenia solution, using copper powder as a catalyst. By using the exidising medium in large excess, he obtained good yields of eyanic acid from methanel (34 g. per 100 g), methyl iodide (22 g), and pinene (16 g.), and even greater yields from more complex acids and

their derivatives. The production of cyanates by the reaction of carbon diexide and ammonia with alkalis such as caustic soda, sedium carbonate, er even sedium sulphide is frequently mentioned in the recent patent literature. The pressure synthesis of urea frem carbon dioxide and ammenia.

CO.+2NH,=CO(NH.).+H.O.

is now an established technical process, and frem the tautement relationship of urea to ammonium cyanate,

CD(NH,), => NH,CNO,

it is not surprising that, if the synthesis is carried out in the presence of alkalis at suitable temperatures, alkalı cyanates result. Such a process is described in B.P. 359559, 1930, the working conditions being 50 atmospheres pressure and a temperature of 400°C. H. Wittek (B.P. 358745, 1930) protects the reaction between ammonia-carbon dioxide mixtures and alkah oxides in the presence of catalysts such as copper, manganese, alumina, and charcoal, and in F.P. 731167, 1932, a heneficial effect is claimed for the addition to the reaction mixture of aubstances which will diminish or prevent the presence of water (which is formed in the reaction), such as earbon monoxide which reacts with it by the water gas reaction. reaction temperature is generally between 500° and 800°C. The technical interest in these processes hes in the possibility of their being utilised as a step towards cyanide production in B.P. 359559, for example, it is stated that cyanide results if the reaction product is subsequently heated with hydrogen, and in BP. 358745 the addition of earhon to the solid raw material is proposed for the same purposa.

material is proposed for the same purpose. Cyanic acid, HCNO, is prepared by heating its polymer cyanura acid, H₃C₃N₃O₃, which is itself obtained by the action of water on cyanuria bromide, C₃N₃Br₃, the triple polymer of cyanogan hremide (p. 504). It may also be compared to the compared prepared by warming urea with phosphorus pentoxide (Weltzien, Annalen, 1858, 107, 219), and condensing the evolved vapour at -18°C. It is a volatile liquid, having a smell resembling that of atrong acetic acid. It is extremely unstable and changes repudly, even at 0°C. that they polymerise to trimeric cyanurates as into the white amorphous polymeric substance cyamelide. Above 0°C, the change is explosively violent, and cyanurie acid is also formed. The by the action of alkalis on cyanogen (p. 503), added, the triple polymer cyanure sed at or en cyanogen braide (p. 504), and by the formed. The relationship between cyanic seld, oxidation of cyanides on heating an are (p. 489, HCNO, cyanorico cid, (HCNO), and cyanody.

studied by Troost and Hautefeuille (Compt.) rend. 1868, 67, 1340) and by van't Hoff and van Deventer ("Studies in Chem. Dynamics," 1896). All of these substances yield the same vapour: at 150°C, and 50 mm, there is a triple point at which the two solid phases cyamelide and cyanuric acid co-exist together with the vapour phase. Above 150°C. cyanuric acid is the stable solid phase, and below 150°C. cyamelide. Cyanic acid is quite unstable (having a higher vapour pressure at the same temperature) with respect to cyamelide.

The aqueous solution of cyanie acid is very unstable, rapid hydrolysis to ammonia and carbon dioxide occurring on warming:

$HCNO+H_2O=NH_3+CO_2$

Ammonium cyanate, NH4CNO, is prepared as a white snow by mixing the vapour of cyanie acid, well diluted with an inert gas, and gaseous ammonia. On warming the solid, or its aqueous solution, ammonium cyanate is converted into urea: NH₄CNO=CO(NH₂)₂. The reversibility of the conversion and its equilibrium in aqueous solution have been investigated by Walker and Hambly (J.C.S. 1895, 67, 746); at 100°C., in N/10 solution, 5% of the eyanate remains unconverted at equilibrium, and the equilibrium is little affected by temperature.

According to B.P. 354604, 1930, ammonium cyanate may be used as a starting point for the preparation of the cyanates of the metals of the first two groups of the periodic table, by treating it in alcoholic solution or liquid ammonia solution at 30-35°C. with the appropriate metallic oxide

or hydroxide.

Potassium cyanate, KCNO, is readily prepared by grinding together dry potassium cyanide and the theoretical quantity of lead oxide, and heating the mixture to the meltingpoint of lead:

KCN+PbO=KCNO+Pb

The salt is recrystallised from water or dilute alcohol. J. Milbauer (Z. anorg. Chem. 1904, 42, 433) observed the conversion of thiocyanate into eyanate by the agency of lead oxide at 300°C.:

KCNS+PbO=KCNO+PbS

The oxidation of cyanide may also be carried out electrolytically, by electrolysing a solution containing 4-6 mols. of KCN and 1 mol. of KOH per litre at 4-6 volts, and a current density of 1-4 amps. per square decimetre of anode surface, agitating the liquor at the anode (Paternò and Pannain, Gazzetta, 1904, 34, (ii), 152). The oxidation is almost quantita-

Potassium cyanate crystallises in thin transparent plates of sp.gr. 2.05, readily soluble in water and anhydrous ammonia, insoluble in absolute alcohol. It is fusible below red heat without decomposition.

Sodium cyanate, NaCNO, may be prepared by similar methods to those described under potassium cyanate. Its solubility in water is 10.68 g. per 100 g. of water at 16°C. It fuses without decomposition at about 500°C.

Lead cyanate is a crystalline precipitate formed when solutions of a lead salt and a soluble eyanate are mixed. The dry salt is stable, but hydrolyses on boiling with water into urea and lead carbonate.

Silver cyanate is slightly soluble in cold water (0.006 g. per 100 c.c. at 12°C.), but much more soluble in hot water. It is readily soluble in aqueous ammonia, and sparingly soluble in

cold dilute nitric acid.

VIII. THIOCYANATES.

The thiocyanates or sulphoeyanides, the sulphur analogues of the cyanates, are not in great demand in commerce, and the fact that some synthetic processes for their manufacture have at times excited technical interest is due to the possibility of converting them into the valuable cyanides (see Alkali Metal Cyanides, p. 480). The commercial requirements of thiocyanates are at present completely satisfied by the gas works production, and this source could easily be expanded to meet any possible increase in the demand.

TECHNICAL METHODS OF PREPARATION.—
1. From Ammonia and Carbon Disulphide. This synthesis was first brought into practical operation by Gelis (B.P. 1816 of 1860; see also Hofmann, Reports Juries, Intern. Exhib. of London 1862, 59). A concentrated solution of ammonia and ammonium sulphide was agitated with carbon disulphide, yielding a solution of

ammonium thiocarbonate,

$$(NH_4)_2S+CS_2=(NH_4)_2CS_3$$

which was then heated to 90-100°C, with potassium sulphide:

$$2(NH_4)_2CS_3+K_2S$$

=2KCNS+2NH₄SH+3H₂S

Günzberg and Tcherniae (B.P. 1148 of 1878, 1359 of 1879, 1261 of 1881) avoided the troublesome formation of large quantities of sulpluretted hydrogen by heating 20% ammonia solution with carbon disulphide at 100°C. under pressure in stirred vessels. The reaction in this case is (Conroy, J.S.C.I. 1896, 15, 8):

$$4NH_3+CS_2=NH_4CNS+(NH_4)_2S$$

The formation of ammonium sulphide increases the ammonia consumption and is responsible for the pressure developed during the reaction. Crowther and Rossiter (B.P. '7846 of 1893), and Brock, Hetherington, Hurter and Raschen (B.P. 21451 of 1893) overcame both drawbacks by adding lime to the charge, when calcium thioeyanate and hydrosulphide are formed:

$$2NH_3+2CS_2+2Ca(OH)_2$$

= $Ca(CNS)_2+Ca(SH)_2+4H_2O$

To avoid the formation of calcium thiocarbonate, the reaction must be carried out in the presence of an excess of ammonia, which is afterwards distilled off. The residual mixture is treated with carbon dioxide to remove sulphuretted hydrogen and to precipitate the equivalent calcium carbonate, and the clear solution of calcium thiocyanate used for the preparation of with the ammonium cyanide present to give the thiocyanates of other metals.

 From Cyanamide and Sulphur.—A process is disclosed in B.P. 384662, 1931, and 412254. tbiourea is converted into thiocyanate and ammonia. If salts of the alkali metals are present, e g. potassium sulphate, a solution of potassium thiocyanate can be obtained by leaching the reaction product. The temperature at which cyanamide can be converted into thiocyanate is thus much lower than that at which it can be converted into cyanide (see Alkaline Earth Cyanidea, p. 488), but considering the thiocyanste process as a step in a evanide synthesis, it is doubtful whether this advantage of lower working temperatures can compensate for the fresbly precipitated cuprous oxido (B.P. 11984 further complications involved in ronverting the thiocyanate into cyanide (see Alkah Metal Cyanides, p. 480).

3. From Coal Distillation Gases .- Coal distilla tion gases are at present the sole source of thiocyanates, which are found in spent oxide from

sumonia still efficent.

SPENT OXIDE may contain 0-10% of summonium thiocyanate, depending on the composition of the gas being purified and on the conditions under which the exide mass has been regenerated (rapid action with consequent rise of temperature favours thiocyanate formation). The whole of the cyanogen contained in the spent oxide can be recovered as calcium thiooyanate by hesting with milk of hime under pressure above 100°C. (Marasae, G.P. 28137), when the iron cyanide compounds and the free aulphur in the mass react to give calcium ferrocyanide and calcium sulnhite, which then react together with more sulphur to give calcium thiooyanate and calcium sulpbate; the iron appears as ferrous aulphide. Holbling (Z. angew. Chem. 1897, 10, 162, 297) the reaction is only 70% complete, but the conversion is improved by using baryta or barium aulphide instead of lime, when a maxture of harium thiocyanate and harium thiosulphate is produced.

Gas Liquon is virtually a very dilute solution of thiocyanate, as ia shown by the following figures relating to Linder'a analyses of five English gas liquors (Chief Inspector of Alkalı

Works, 42nd Rep., 1905, 35):

Analysis in g /litre. Sample. HCN as ferrocyanide 0.068 0.0 0 576 0-460 0 0 HCN as thio. cvanate 1.688 4.457 1.238 1 029 1 042 HCN as bydroevanio acid. 0 0 0 0 8 3 0 0 030 0 050 0 355

thiocyanate.

hydrogen and some hydrocyanic acid) evolved are passed into sulphuric acid for the production of ammonium aulphate. The waste liquor from the stills contsum about 0 15% of thiocyanate in the form of the calcium salt, which is best recovered as the cuprous salt by tresting the acidified hiquor with copper aulphate and a reducing agent such as sulphur dioxide (Parker and Robinson, B.P. 2383 of 1889), or by treating the liquor (after removing excess of calcium hydroxide with carbon dioxide) with of 1893).

FORMATION OF THIOCYANATE DIRECTLY FROM COAL GAS -The hydrocyanio acid present in coal gas can be quantitatively fixed as thio cyanate under suitable conditions. necessary sulphur for the reaction can be gas purification, and also in gas liquor and provided by acrubbing the gas with a sodium olysulphide solution (Marriott, B.P. 3311 of 1877), or with a auspension of manganese dioxide in water, which produces aulphur by oxidising sulphuretted hydrogen in the gas (Campbell and Boyd, B.P. 10186 of 1888) H. Wood Smith, Gidden, Salamon and Albright (B.P. 13653 of 1901) proposed simply to scruh the gas with an aqueous suspension of aulphur, obtaining a solution containing up to 200 g. of ammonium thiocyanate per litre. This process is in operation at several gas works. In the process described by A. E. Broadburry (J. Gas Lighting, 1012, 31; B.P. 23624 of 1909) the aulphur is provided by spent oxide. which is charged into an ordinary purifier box and continually aprayed with water to keep it According to moist, while the gases pass upwards through it. The reactions are:

$$H_2S+2NH_s+S=\{NH_s\}_2S_2$$

 $\{NH_s\}_sS_s+HCN+NH_s$
 $=\{NH_s\}_sS+NH_sCNS\}_s$

In order to prevent water-logging, the direction of the gas flow is changed at intervals, and the apent oxide charge is removed after 6-8 weeks The capacity of a 20 ft. hy 20 ft. purifier box containing a 2 ft. layer of spent oxide is 3 5×10° cu. ft. of gas per day, and 3-5 lb. of ammonium thiocyanate are obtained per ton of coal, in the form of a 26% solution.

PROPERTIES OF THE THIOCYANATES.-Ammonium thiocyanate, NH, CNS, crystalluca in anhydrous, colourless leaflets of apgr. 1-3 and melting point 159°C. 100 g. of water dissolve 122 g. of the salt at 0°C, and 162 g. at 20°C. It is readily soluble in alcohol. The fused salt The formation of ammonium thiocyanate in the isomeric thiourea. At 220°C, decomposition gas liquor is due to the action of dissalved air on into the volatile products ammonia, sulphuretted the ammonium sulphide in the fresh liquor, hydrogen and carbon bisulphide occurs, and giving ammonium polysulphides which react guandine thiocyanate remains behind.

Potassium thiocyanate, KCNS, crystallises in anhydrous, striated prisms or needles insecticides (U.S.P. 1841458, 1932; Murphy of sp.gr. 1.9 and melting-point 172.3°C. It is and Peet, J. Econ. Entomol. 1932, 25, 123; deliquescent and very soluble in water:

25 Temp. (°C.) 20 177.2217 239 Solubility (g./100 g.)

It is also soluble in anhydrous ammonia, and in various organic liquids as follows:

Solubilities (g./100 g. of solvent). 20.4 at 58° Acetone 20.8 at 22° Ethyl

acetate 0.44 at 0° 0.2 at 79° 0.4 at 14° Amyl

2·14 at 100° alcohol 0.18 at 13° 3.21 at 115° 6.75 at 0° 4.97 at 58° Pyridine

The fused salt becomes deep blue at 430°C., probably due to the liberation of sulphur.

Thiocyanogen, (CNS)2, is obtained by the action of a solution of a halogen in an anhydrous solvent on a metallic thiocyanate; for instance, from a dry ethereal solution of bromine cooled to 0°C. and lead thiocyanate. On cooling an approximately N/2 solution of thiocyanogen in carbon bisulphide to -70° C., almost colourless crystals melting at -3° to -2° C. are obtained, which decompose at the ordinary temperature. The chemical behaviour of thiocyanogen is similar to that of the halogens: its electronegative character is at least equal to that of iodine. The molecule is believed to contain two thiocyanogen radicles, and to have the constitution N:C·S·S·C:N (Soderback, Annalcn, 1919, 419, 217). A standard solution is used for determining the thiocyanogen values of fats; with compounds containing one double bond they resemble the iodine numbers, but only one double bond in linoleic acid is attacked. The method has been used for finding the composition of natural oils (Freudenberg, Arch. Pharm. 1925, 263, 675; Z. Unters Lebensm. 1926, 51,

USES OF THIOCYANATES.—Thiocyanates do not find large-scale application in technical industry, the annual world output being estimated at 150 tons (Chem. Trade J. 1936, 98, 434). The aluminium and tin salts have been used in the dyeing and printing industry to a small extent. and the yellow compound canarin, (q.v.) obtained by oxidising potassium thiocyanate with a strong oxidising agent such as chlorate and hydrochloric acid, has been used as a yellow dye for wool. Ammonium thiocyanate is used as a fixing agent in photography.

Thiocyanates are plant poisons, and their use (especially in the form of the ammonium salt) as weed-killers has been proposed (see Chem. Trade J. 1936, 98, 498). Frankland and Silvester (J.S.C.I. 1907, 26, 231) investigated the germicidal properties of thiocyanates. They found that goldfish were not affected in 24 hours by concentrations of up to 260 parts per 100,000 of thiocyanate in the form of the potassium salt, but that as little as 10 parts per 100,000 were sufficient to control Bacillus coli communis. Thiocyanates are present in the animal organism, in the human saliva, gastric juice, conjunctiva secretion and urine.

Organic thiocyanates have been proposed as Hartzell and Wilcoxon, B. 1936, 612).

Analytical.

The methods of analysis described here are those commonly employed in technical laboratories. For further details, an analytical textbook or a work on technical analysis should be consulted.

The Detection of Cyanide Compounds is based chiefly on two characteristic, deeply coloured cyanide derivatives, namely Prussian blue and the red ferric thiocyanate. The Prussian blue reaction is given directly by ferrocyanides when their neutral or acid solutions are treated with ferric salt solutions, a blue precipitate or a blue coloration due to ferric ferrocyanide being produced. The reaction may be applied to simple cyanide solutions (after evaporating them down to small bulk with a little alkali if they are very dilute) by treating them with ferrous sulphate solution to form ferrocyanide, and then producing Prussian blue by adding ferric salt solution, or better by adding an oxidising agent such as nitric acid to convert the excess of ferrous salt into ferric salt, with the same result. The latter procedure is stated to give quicker pre-cipitation of Prussian blue, and greater sen-sitivity (Vorlander, Ber. 1913, 46, 181). Verhoever and Johns (J. Amer. Chem. Soc. 1915, 37, 601) give details of a method of this kind. The dilute cyanide solution is evaporated, after the addition of a little sodium hydroxide, in vacuo at a temperature below 70°C. to a volume of I ml., 0.2 ml. of 3% FeSO₄ and 0.05 g. of KF are added, and after 10 minutes a small excess of 30% HNO₃. The blue colour is compared with that produced by a known quantity of KCN under similar conditions. For a similar method Kolthoff (Z. anal. Chem. 1919, 57, I) claims a sensitivity of 2 mg. CN per litre.

A deep red coloration due to ferric thiocyanate is produced when a ferric salt is added to a neutral or acid thiocyanate solution. In the case of very dilute solutions, it is advantageous to shake the treated solution with a small quantity of ether to extract the ferric thiocyanate from the water. This test can be applied to cyanides by converting them into thiocyanates by means of ammonium polysulphide solution and can be made quantitative by matching the coloration produced with that of a standard thiocyanate solution. Johnson's method (J. Amer. Chem. Soc. 1916, 38, 1230) is as follows: an HCN solution containing 0·1-8 mg. of HCN is made alkaline with KOH, treated with I ml. of ammonium polysulphidesolution, and evaporated to dryness; the residue is extracted with acetone, which dissolves only the KCNS, and the acetone removed by evaporation; the residue is dissolved in water and treated with 2 ml. of 0.5% FeCl₃, the colour being matched by means of a standard thiocyanate solution. Castiglioni (Gazzetta, 1933, 63, 171) treats the aqueous cyanide solution with sulphur dissolved in acetone and claims a sensitivity of I in 50,000.

Another sensitive colour test for cyanides is are titrated with pure silver nitrate solution the sodium picrate test, which gives a reddishviolet coloration depending on the formation of sodium isopurpurate when the cyanide is in excess, and of amino-nitrophenols when picrate is in excess (A. C. Chapman, Analyst, 1910, 35, 469). The liquid to be tested is mixed with an equal volume of a solution containing 0 05% pieric acid and 0 5% Na₂CO₃, and digested at 40°C, for I hour. The colour is compared with that produced by a standard HCN solution, and the sensitivity is 1 part of HCN in 500,000 of water (Waller, Proc. Roy. Soc. 1910, 82, B,

Several colour tests have been proposed for the detection of minute concentrations of HCN in the atmosphere of ships and buildings after fumigation. In the benzidine copper acetate test (Pertusi and Gastaldi, Chem -Ztg 1913, 37, 609; Hamer, J. Roy. Sanit. Inst 1933, 53, 563), the cyanogen formed by the action of HCN on cupric acetate oxidises benzidine to a blue substance. The test is carried out by dipping a filter paper into a mixture of equal volumes of a 0.3% solution of copper acctate and a 0.1% solution of benzidine acetate, draming surplus eclution from the paper and introducing the damp paper into the atmosphere to be tested HCN concentrations of 0 002% and higher are readily detected, a blue coloration being developed in 10 seconds, but the rapid action makes it difficult to cetimate the actual concentration. In the mercure chloride test, the action of HCN on HgCi, to give non ionised Hg(CN), and HCl is detected by having present an indicator with a well marked colour change in the appropriate $p_{\rm H}$ range, e.g. methyl orange (Sberrard, Fublie Health Reporte, Reprint No. 1224, 1228, Treasury Dept, U.S. Public Health Service). The methyl orange mercuno chloride test is much slower than the benzidine copper acetate test, and is therefore better adapted to the estimation of actual HCN concentrations.

Cyanates are best characterised by the deep hine potassium cobalt cyanate, obtained by is by precipitating PbS by means of lead adding potassium and cobalt salt solutions to carbonate and cotadieng the precipitate with the cyanate solution made and with action boiling mitric acid to PbSO, which is washed acid. Cyanide must be removed before the and weighed.

cobalt solution is added. The nitroprussides give a deep violet coloration with alksh sulphides. The colour is obscured if heavy metals which form dark-coloured sulphides have not been previously

removed from the solution.

Analysis of Alkals Cyanides.-Sodium and potassium cyanides are hygroscopic when cold, so that care is necessary in taking samples for analysis. The samples should be transferred immediately and without touching them with the hands to a hot iron mortar, coarsely crushed, and transferred to a well dried bot bottle. preferably fitted with a lightly greased glass stopper. Errors due to absorption of moisture in sampling occur very easily, and may be more serious than inaccuracies in testing

The Estimation of Cyanogen is carried out by a modified Liebig's method. 5 g. of the sample are weighed out and made up to 500 ml, and 25 excess of Ba(NOs), is then added to pre-

until a faint turbidity is visible. The reaction is

2NaCN+AgNO3=NaNO3+NaAg(CN)... and when this is completed, the further addition of silver nitrate leads to the precipitation of silver cyamde, thus:

NaAg(CN)2+AgNO3=NaNO3+2AgCN

The eilver nitrate solution must be neutral and free from lead and zinc. It is standardised by means of a solution of pure sodium chloride (dried by heating to redness before weighing), using pure potassium chromate as indicator In impure solutions of cyanides, the end point is more easily observed if a little caustic sods and a few drops of 10% K1 solution are added before running in the silver nitrate solution. Wellings (Analyst, 1933, 58, 331) suggests the use of dipbenyl carbazide as an absorption indicator. the end point being marked by a colour change from pink to violet Sulphides interfere with the test, but they are easily removed by shaking the solution with a little lead carbonate and filtration. The usual strength of eilver nitrate solution is N/10, of which I ml is equivalent to 0 0098 g NaCN For very dilute cyanide solutions, N/100 silver nitrate may be used,

Sulphide is generally estimated by Ewan'a method (JSCI 1909, 28, 10; see also Rossiter, JSCI 1911, 30, 583) 10 g of the finely powdered sample are dissolved in 15 ml of ordinary water and a lead nitrate solution containing 2 4 g per litre is run in gradually. The mixture is stirred to coagulate the colloidal lead sulphide into brown flocks, and the addition of lead natrate continued until a drop of the liquid placed on a filter paper produces no dark coloration with a drop of lead nitrate solution The results obtained are low, but for small sulphide contents are made sufficiently accurate by taking I mol. of the standard lead nitrate solution as equivalent to 0 0007 g Na,S (i.e. by multiplying the theoretical result by 125) The alternative method of sulphide estimation

Water is estimated by weighing out about I g of the coarsely powdered sample into a small distilling flask (10-20 ml), which is then evacuated by a good water pump or hy a mercury pump. The flask is then gently warmed, finally to 200-300°C., when the whole of the moisture is expelled in a few minutes without appreciable decomposition of the cyanide.

Carbonate is conveniently estimated by precapitating a suitable portion of the solution with barium chloride in the cold (to avoid decomposition of any cyanate which may be present). The precipitate is washed and titrated, together with the paper, with hydrochloric acid and methyl orange.

Hydroxide may be estimated by a modification of Clennell'e method (Chem. News, 1895, 71, 93) To 100 ml of the solution, standard AgNO, solution is added to give a permanent turbidity, ml. of this colution, diluted to 200-300 ml, cipitate the carbonate, which is filtered off and filtrate is then titrated with standard acid and and

phenolphthalein.

The Estimation of Chloride is carried out after removing cyanide from the solution by neutralising it with dilute nitric acid, using methyl orange, and boiling for about 1 hour in a draught cupboard. This treatment also decomposes any cyanate present. The chloride remaining is then precipitated by a known excess of standard AgNO_s solution, the AgCI filtered off and the excess of silver back-titrated with ammonium thiocyanate and iron alum, in the usual way. If ferrocyanide is present, it may be precipitated with excess of $AgNO_3$ in ammoniacal solution; on acidifying the filtrate with nitric acid, AgCl is precipitated and may be weighed.

Cyanamide in the presence of alkali cyanides is readily estimated by exactly neutralising the solution of 1 g. of the substance in 100-150 ml. of water with nitric acid, and boiling for 20 minutes to expel HCN. After cooling, a few drops of ammonia and then an excess of AgNO₃ are added. The canary-yellow precipitate of silver cyanamide is filtered off and dissolved in cold dilute nitric acid, when a small residue of undissolved AgCN is usually left. The silver in the solution is then titrated against ammonium thiocyanate with ferric alum indicator.

This method may be used for cyanamide in the absence of cyanide, when the expulsion of HCN by means of nitric acid may be omitted. For substances rich in cyanamide, it is better to estimate the nitrogen content of the silver cyanamide precipitate by Kjeldahl's method, for although it contains all the cyanamide nitrogen, the precipitate in this case is not pure Ag₂CN₂ (Caro, Z. angew. Chem. 1910, 23, 2405).

Cyanate is estimated by acidifying the solution with nitric acid and boiling for at least 20 minutes in a draught cupboard. The cyanide is thereby expelled as HCN, and the cyanate is converted into ammonium nitrate and carbon dioxide:

HCNO+HNO3+H2O=NH4NOS+CO2

After cooling, the solution is boiled with excess of caustic soda and the ammonia evolved is absorbed in a known excess of standard acid and estimated in the usual way. If cyanamide is present, some ammonia originates from it during the boiling with caustic soda, and in this case Ewan (J.S.C.I. 1904, 23, 244) suggests that the CO₂ evolved when the sample is boiled with acid should be collected and measured in order to estimate the cyanate.

Analysis of Ferrocyanides.-The most convenient method of ferrocyanide estimation is by titration with a standard zinc sulphate solution. precipitate containing zinc ferrocyanide is obtained, and the titration is continued until a drop of the solution no longer gives a blue colour with a drop of ferric solution. The procedure is as follows: a solution containing 10.2 g. of $ZnSO_4.7H_2O$ and 10 ml. of concentrated sulphuric acid per litre is standardised against a solution of pure potassium ferrocyanide (10 g. K_4 Fe(CN)₆:3H₂O per litre), 25 ml. of which is mixed with 25 ml. of a saturated solution of K_2SO_4 , 25 ml. of water and 10 ml. of 10%

washed with CO2 free distilled water, the sulphuric acid. Equal volumes of the standard washings being added to the filtrate. The zinc and ferrocyanide solutions are required, and the precipitate has the composition $Zn_3K_2[Fe(CN)_6]_2$ (Meurice, Ann. Chim. Analyt. 1913, 18, 342). Subsequent titrations should be made as nearly as possible under the same conditions. The addition of an excess of potassium sulphate is made with the object of eliminating variations in the composition of the precipitate and thus in the quantity of zinc solution required, depending on the nature of the metal in the ferrocyanide used (Colman, Analyst, 1908, 33, 261; J.S.C.I. 1908, 27, 806; see also Treadwell and Chervet, Helv. Chim. Acta, 1923, 6, 559).

Feld's mercuric cyanide method (J. für Gasbeleuchtung, 1903, 46, 565; J.S.C.I. 1903, 1068) is more accurate. Ferrocyanide equivalent to 0.3-0.5 g. of yellow prussiate is boiled with 10 ml. of N-NaOH and 15 ml. of 3 M.MgCl₂ solution (285 g. MgCl₂ per litre). The boiling is continued for 10 minutes to expel any cyanide completely as HCN, after which 100 ml. of boiling N/10-HgCl₂ solution are run in, and the boiling continued for 10 minutes to convert the ferrocyanide into mercuric cyanide. The mixture is then distilled for 30 minutes with 30 ml. of 3N-H2SO4, the HCN evolved being absorbed in caustic soda and the cyanide in the resulting solution titrated with AgNO₃, using KI as indicator. Williams (J.S.C.I. 1912, 31, 468) avoids the conversion of ferrocyanide into mercurio oyanide by making use of the catalytic action of cuprous chloride, 0.05-0.10 g. of the pure compound dissolved in HCl being added to the ferrocyanide MsCl₂ (as above) or with Pb(NO_S)₂, which also removes sulphide. On boiling the mixture for 30 minutes, the CN of the ferrocyanide is quantitatively removed as HCN.

Analysis of Thiocyanates.—Volhard's method,

namely, the precipitation of insoluble silver thiocyanate by the addition of AgNOs to a thiocyanate solution which is neutral or acid with nitric acid, is used. The end point is determined by adding a small amount of a ferric salt to the solution before titrating, the disappearance of the red colour due to ferric thiocyanate indicating that the precipitation of the thiocyanate as silver salt is complete. A quantity of the solution containing about 0.25 g. of thiocyanate is diluted to 100-150 ml. with distilled water, acidified with 10 ml. of dilute HNO₅, and 5 ml. of a 10% Fe(NO₃)₃ solution are added. The mixture is titrated with N/10-AgNO₃ solution. In analysing a gas works product for thiocyanate, ferrocyanide is first removed by slightly acidifying the solution and adding ferric alum. The filtrate is treated with acid sodium sulphite in excess, then with copper sulphate, and is boiled for a few minutes. The precipitate of cuprous thiocyanate, which may be mixed with copper sulphide, is filtered off, washed with boiling water, and digested with excess of caustic soda. Cuprous hydroxide

latter being separated by filtration and titrated by Volhard's method above. Analysis of Gas Works Products.-Cyanide

and sodium thiocyanate solution are formed, the

and ferrocyanide in gas purification masses are ; determined by triturating 30-40 g. of the cyanadin glycosides in nearly two hundred material in a mortar with excess of cold caustic different flowers and fruit by G. and R. soda solntion for several hours. A httle Robinson (Biochem. J. 1931, 25, 1687; 1932. FeSO, may be added to convert cyanide into 28, 1647; 1934, 28, 1712), more recent solution ferrocyanide. The filtrato is treated with 4-5 and analysis of cyanidus prements includes that times its velume of methylated spirit, which precipitates all the sodium ferrocyanide present (Biochem. J. 1930, 24, 753), elderherry skins and but retains carbonyl ferrocyanide (see p. 477), thiocyanate, sulphide, and excess of caustic soda in solution. The precipitate is then treated as described above (Colman, Analyst, 1908, 33, 261; J.S.C.I. 1908, 27, 806; Skurrow, J.S.C.I. 1910, 29, 310).

The small quantities of cyanogen compounds in concentrated ammonia liquor may be estimated by converting them into ferme thiocyanate and measuring the depth of colour hy means of a tintemeter (Spielmann and H. Wood, J.S.C.I. 1919, 38, 43T, 369T). Thecyanato initially present is first estimated by acidifying the diluted sample with H,SO, in the cold, and removing ferrocyanide by means of a small excess of iron alum; a large excess of iron alum is then added to the filtrate, which is examined in the tintometer. Cyanido is next converted into thiceyanate by digestion at 30-35°C with ammonium polysulphide solution, and a second estimation of the depth of coleur is made Thiocarhenste is converted into thiocvanate by digestion at 70-75°C. for 45 minutes with NH;

cyanide and ferrocyanide being unaffected.

The HCN content of coal gas is generally determined by treating the gas with ammonium polysulphide solution, when thioeyanic acid, HCNS, is fermed and can be astimated as thiecyanate (see Voituret, Brennstoff-Chem. 1932, 13, 264; Pieters and Penners, Het Gas, 1932, 52, 382). Alternatively the gas may be washed with a mixture of caustic soda, ferrous sulphate, and water (as in a cyanide washer) to obtain the HCN in the form of ferrocyanule, which can he estimated by the mercurie cyanide method already described An account of different methods for the determination of HCN in coal gas is given by Brender & Brandis and

Bohlken, Fet Gas, 1933, 53, 194.

G. E. W. CYANIDIN is the most common and widely distributed of the authocyan pigments of plants,

and occurs naturally in the form of various glycosides.

It was first isolated by Willstätter and Everest (Annalen, 1913, 401, 189) from the blue cornflower pigment cyanin, and may be obtemed hy hydrolysis of this or any other of its naturally occurring glycosides, or from its methyl ether, peonidin, by demethylation. Glucoside-pigments derived from cyanidin have been replated from, or detected in, the flowers of blue and deep purpls cornflowers, Rosa gallica, peony, ehrysanthemum, dahla (deep red), aster, poppy, tracted with other, and the product crystallised Zinnia elegans, gladiolas (hybrid), Gaillardia from water. Readily soluble in water, sloobel. bicolor, Helenium autumnale, Tulspa Gernersana, acetone or glacial acetic acid, it is insoluble in Tropcolum maius, and in the fruits, or berries, benzens; with soda it gives a yellow coloration, of the sweet cherry, sloe, cranberry, red current, with HCl cyanidia chloride is formed and raspberry and mountain ash. In some cases crystallises out. When decomposed by means they are accompanied by derivatives of other of fused alkali, protocatechuse acid and phloroanthocyanidins (see various papers by Wallstatter glueinol are produced. and his collaborators).

Apart from the qualitative identification of and analysis of cyanidm pigments includes that from the crimsen antirrhinum by Scott-Moncrieff flowering current by Nolan and Casey (Proc. Roy. Irish Acad. 1931, 40B, 56; 1936, 43B, 11. American cranberries by Grove and Robinson (Broehem, J. 1931, 25, 1706), blackberries by Karrer and Pieper (Helv. Chim. Acts, 1930, 13, (Dissertation, Zürich, 1918), shiso leaves by Chika Kuroda and Mizu Wada (Proc. Imp. Acad. Tekyo, 1935, 11, 28), mairs by Sando, Milner, and Sherman (J. Biol. Chem. 1935, 109, 203), and hibsens by Yamamoto and Oshima (Sci. Papers Inst. Phys. Chem. Research, Tokye, 1932, 19, 134).

The natural glycosides of cyanidin already known are the 3-glucosides chrysanthemin, etc., the 3-galactoside idaein the 3-diglucoaide mekoeyanin, the 3-rhamnoglucosides heracyansa, anterrhenia, etc., the 3.5 diglucoside cyansa, the 3-pentoside historia, and the hi-

cyamin, the 3-pentosics houses, and the his melecular glucoside and pentoside sambus-cyamin. Acylated glycosides also occur. Cyanidin. Chloride. C.3-fij.O.Cl. 357. trhydroxy-2-mp-dibydroxyphayi.1.4-henco pyrand-anhydrochlerde crystaliaes readly from 20% HOI when its pure glicosides are hydrolysed by holing with this reapon for 3 mantles; the crystalis, long red-movem needle with malable listic, centain 147.57, which is very difficult to remove, complets drying being only obtained at 105°C, in high vacuum. The anhydrous salt does not meit below 300°C; if dried at 50°C, it melts at once if dipped into a bath at 220°C, but does not melt if put in st 200°C, and the temperature then gradually raised. It is very soluble in methyl er sthyl alcohol; fairly soluble in dilute sulphune and; difficultly soluble in HCl. Neutral alcoholic or aqueous solutions lose their colour en standing owing to pseudo hase formation, acids reproduce coleur, alowly if celd, rapidly if warm. The absorption spectrum shows one broad hand with all defined edges.

Cyanidin colour base separates in crystallino condition when a hot concentrated solution of the chlorids in alcohol is mixed with twice its volume of water (cf. Willstätter and Nolan, Annalen, 1915, 408, 13). It is fairly soluble

m alcohol er pyridins.

Cyanidin pseudo base, C₁₅H₁₂O₇, crystal lises with 1H₂O when a dilute solution of the chlorele in alcohol is mixed with twice its volums of water, warmed gently, then, after it has become colourless, evaporated in vacuo, ex-

Cyanidm chloride dyes wool (tin mordant)

blue-violet, cotton (tannined) violet, and unmordanted wool fine rose (Willstätter and Mallison).

Qualitative analytical tests for cyanidin include the reddish-violet colour of its amyl alcohol solutions on shaking with sodium acetate, turning to a pure blue with a drop of ferric chloride, the incomplete extraction hy the "cyanidin reagent" and "delphinidin reagent," the moderate resistance to oxidation in alkaline solution, and the complete precipitation from amyl alcohol hy 5-6 volumes of benzene. Cyanidin can also he identified by the specific colour reactions of its various glycosides (G. Robinson and R. Robinson, ibid.).

Many different syntheses of cyanidin have now been carried out, and these confirm the structure of the chloride which was first put

forward hy Willstätter and Everest. The claims of Nierenstein and Malkin (Ber. 1928, 61, B, 791; J. Amer. Chem. Soc. 1930, 52, 2864) that natural cyanidin chloride is not correctly represented by this accepted formula, have heen dealt with hy Rohinson and Rohertson (J.C.S. 1928, 1503), Rohinson (Chem. Age, 1928, 19, 337), Willstätter and Rohinson (Ber. 1928, 61, [B], 2504), Kuhn-Jauregg (ibid. 2506).

The first synthesis was carried out by reduction of the related flavonol quercetin hy Everest (Proc. Roy. Soc. 1914, 87, B, 444; 1914, 88, B, 326) and by Willstätter and Mallison (Sitzungsher. Preuss. Akad. Wiss. Berlin, 1914, 769). More recently Asahina and Inubuse, using sodium amalgam in alkaline solution, have obtained cyanidin by the reduction of rutin (3-rhamnoglucoside of quercetin) and also, with more difficulty, of quercetin itself (Ber. 1931, 64, B, 1256). Karrer, Yeu, and Reichstein (Helv. Chim. Acta, 1930, 13, 1308) have obtained small yields, using titanium trichloride as a reducing agent. Anthocyanidins have also heen obtained from flavones by Kaoru Kondo and Hisao Segawa (J. Pharm. Soc. Japan, 1932, 52, 353, 358).

Willstätter, Zechmeister and Kindler (Ber. 1924, 57, [B], 1938) synthesised cyanidin chloride using trimethoxycoumarin, and magnesium 3:4-dimethoxyphenyl iodide, and Robinson and his collaborators have also synthesised it hy various methods, the most satisfactory (J.C.S. 1928, 1526) involving the use of O-benzoylphloroglucinaldehyde.

Appel and Robinson (J.C.S. 1935, 426) have obtained pure cyanidin chloride by oxidation of tetramethyl-d-catechin by treatment with

bromine followed by demethylation with hydriodic acid and phenol. Freudenberg, Fikentscher, Harder, and Schmidt (Annalen, 1925, 444, 135), on the other hand, have obtained small yields of dl-epicatechin from evanidin.

The investigations by G. and R. Robinson (Biochem. J. 1933, 27, 206; 1934, 28, 1712) on the existence and occurrence of leuco-anthocyanins show that the greater proportion give cyanidin chloride on treatment with hot hydrochloric acid.

In the case of such natural substances as the anthoxanthins (flavones and flavonols), tannins, catechins, and leuco-anthocyanins, as well as the anthocyanins, the 3':4'-dihydroxy grouping is by far the most common, a fact which affords a more reasonable explanation for the co-production of glycosides of cyanidin and the related flavonol quercetin in rose, wallflower and maize than the suggestion of Everest (Proc. Roy. Soc. B, 1918, 90, 251) and Sando (l.c.) that the cyanidin in these plants is derived directly from quercetin. The inverse correlation in the production of cyanidin and the unrelated flavone apigenin in Antirrhinum (Scott-Moncrieff, I.c.) and Dahlia (Lawrence and Scott-Moncrieff, J. Genetics, 1935, 30, 155) points rather to the metaholism of anthocyanins and anthoxanthins by parallel syntheses, each pigment being partially derived from a common component whose availability is strictly limited (Scott-Moncrieff, ibid. 1936, 32, 117).

The genetical relationship hetween cyanidin pigmentations and those by the more oxidised delphinidin and the less oxidised pelargonidin is such that, when general anthocyanin production occurs, the more oxidised type is dominant to the less oxidised, and is also deeper and more hlue-toned, e.g. in sweet pea the purple flower contains the delphinidin, the paler red the cyanidin, and the even paler salmon the pelargonidin type of anthocyanin (Scott-Moncrieff, ibid. 1936, 32, 117).

R. S.-M. CYANIN, the anthocyan pigment of hlue or purple cornflowers, also occurs in Rosa gallica, deep red dahlias, hrown wallflowers, etc. It is a diglucoside of cyanidin yielding cyanidin (1 mol.) and glucose (2 mols.) on hydrolysis. Willstätter and Everest (Annalen, 1913, 401, 189) first obtained it in a pure condition, as the crystalline chloride, from the blue cornflower; it is, however, more readily prepared from Rosa gallica (Willstätter and Nolan, Annalen, 1915, 408, 1), or the dcep red dahlia (Willstätter and Mallison, 1915, 147). The Annalen, 408, C27H31O16Cl, red-hrown rhombic leaflets containing $2\frac{1}{2}$ mols. of water, can only be completely dried at 105°C. in high vacuum. Air-dried it is very difficultly soluble in water, acetone or chloroform, difficultly soluble in cold alcohol or dilute HCl, insoluble in benzene. In 7% sulphuric acid it is fairly soluble, but the sulphate separates out from the solution. Soda added to an acid solution gives a pure blue and sodium acetate a violet colour; ferric chloride gives a fine blue in alcohol, violet in water. It shows a single absorption band covering the green and blue; it is optically active. Willstätter and

514 CYANIN.

Nolan give, for white hight, [a] — 25% (±107). The low "distribution number "(partition value between am; la licohol and aqueous HCl) is normal for a digylecoade Perrate, red needs, very coluble in water. Cycans polarsus sell, the blue pigment of the comflower, was obtained crystaline, but not free from NaCl, by Willister and Severes, after purification by dadysas It is very soluble in water, insoluble in atcohol; It is very soluble in water, insoluble in atcohol; a standing owns; to formation of a possess, one whereas in 20%, NaCl solution the colour is stable for months.

Cyann was the first anthocyan to be isolated [828] and euccessfully analyzed It is very wately datributed G. and R. Robinson (Biochem. J. 1931, 25, 1637, 1934, 25, 1647, 1934, 25, "mn lundreds of flowers and fruits that cyann with its dibydroxy grouping is by far the most common type, as is also the case with other prep flavan eutletances Cyann has also been lasolated from brown wallflowers by Scott 1873 Monereff (Chem Ind. Rev. 1929, 7, 630) and from chiso leaves by Chika Kuroda and Mazio free Watel (Proc. Imp. Aced Tokyo, 1935, 11, 28), aw ho found that here it was sometimes combined essays

with p coumane acid.

The position of the glucose molecules was uncertain for some time, Richmon favouring the 6 pointion and Karret the 3. These apparently opposing views have been unted by Robinson's recognition that eyanin, pelargonin, peonin, makin, petum and delpha are all 3.5 dimoso sides (JCS 1931, 2665). This was confirmed by the synthesis of 3.5 & digitactively eyargandic chloride (III) by Todd and Robinson (JCS, 1932, 2693) by condensation of 2.0-mono acetylglucoudylphloroglucinaldehyde (I) and co-0 tetra acetyl & glucosodicy.34 disacetory acetophenone (II), and subsequent deacetylation with KOH.

The 5-monoglucoside, cyanenin (IV), which is not known to occur naturally, can be obtained from cyanin by partial hydrolysis and has also been synthesised by Léon and Rohinson (J.C.S. 1932, 2221).

Since the pg of the cell cap of blue cornflowers is slightly more acid than that of the red Ross Galling, the blueness of the former may be dependent upon a colloidal phenomenon (G. and R. Robinson, Nature, 1933, 132, 525)

R. S. M

CYANINE BLUE, Letch's blue, A "mixed" pigment of cobalt blue and Prussian blue.

CYANINE DYES. The first eyamne dye, prepared in 1856, appeared at the time to be valueless owing to its instability to light. In 1873 Vogel discovered that various dies, amongst them eyanine, possess the power of rendering photographic plates, which normally are sensitive only to light of short wave length, eensitive also to other wave-lengths, the sensitis ing maximum approximating to the absorption maximum of the dye (Ber. 1875, 8, 1635) Whilst very interesting scientifically, even this action of cyamine science at the time to bave little practical value, since it was accompanied by veiling and staining of the plate. In 1903, however, it was found that the related tsocyanines censitise, without accompanying defects, and a flood of German patents was the result, whilst the German chemists arrived at the chemical atructures of cyanine and isocyanine. At the time of the Creat War, it was processary that the Albed Armes should have photographic sensitisers, because they are essential for aeroplane photography. This led Mills to has study of the cyannes, and he and his pupils established the structure of the whole group; thue Pinacyanol, then the best consituer for red light, had been discovered in 1905, but its constitution was not elucidated until 1920, and Thuzele Purple, discovered in 1887, was of unknown etructure until Mills's publication in 1922. The preparation of cyanines containing nuclei other than the original one, quinoline, greatly enlarged the field, as did observations that the new types of eyamnes also had valuable photographic properties; some of them are greatly superior to the older types

At present certain cyanines, absorbing light of comparatively short wave-length, are used for conferring extra sensitivity upon gelatinechlordie photographic emulsions, such as are used for coating gaslight printing papers; others, absorbing longer wave lengths, are used to sensitise gelatino-bromide photographic coulsions, such as are used for plates and him, to any desired part, or to the whole, and the servicine in the control of the properties of the photographic material, which is useful for distance photography and for certain kinds of distance photography and for certain kinds of sextentific work. The cyanine dyes have not only proved valuable to astronomers and other which are accepted to-day, the former with a scientists, but also to the world at large, for 4:4'-linking, the latter with a 2:4'-linking their power of conferring sensitivity to light of various colours has made possible a true rendering of colour in monochrome and led to a great increase of speed, especially to artificial light. The high achievements of modern photography are in fact largely due to photographic sensitisers, of which the cyanines are amongst the most important. Sensitisers also made possible photography liave colour.

Cyanines (4:4'-Cyanines), isoCyanines (2:4'-Cyanines), and ψ-Cyanines (2:2'-Cyanines).— The blue dye, cyanine (Quineline Blue), was prepared by Greville Williams by the action of caustic alkali on the alkiodide of impure quinoline (Trans. Roy. Soc. Edin. 1856, 21, 377; Chem. News, 1859, 1, 15; 1860, 2, 219). Hofmann recognised that this quinoline contained lepidine (Proc. Roy. Soc. 1863, 12, 410), and Hoogewerff and van Dorp showed that the ethiodides of the two bases condense, under the influence of potassium hydroxide in aqueous solution, to form the cyanine molecule according to the equation

$$\begin{array}{l} C_{9}H_{7}N\cdot C_{2}H_{5}I+C_{10}H_{9}N\cdot C_{2}H_{5}I \\ = C_{23}H_{23}N_{2}I+H_{2}+HI \end{array}$$

(Rec. trav. chim. 1883, 2, 28, 41, 317; 1884, 3, 337). They found that a purple dye, isocyanine, is produced by the action of alkali on the alkiodides of quinoline and quinaldine, and they represented its formation similarly (ibid. 1883, 2,317; 1884, 3, 337). Spaltcholz independently discovered the isocyanines, but regarded them as containing two extra hydrogen atoms (Ber. 1883, 16, 1847). He, and also Hoogewerff and van Dorp, pointed out that, in preparing cyanines and isocyanines, it is advantageous to use twice the calculated amount of quinoline alkiodidc. Miethe and Traube's discovery of the value of the isocyanines as photographic sensitisers (G.P. 142926, 1902) led to patents by various German firms, in which substituents were introduced to improve the sensitising action, and the acid radical was altered so as to increase the solubility (G.P. 158078, 167159, 167770, 170048, 170049, 1903). This was largely the work of E. König. isoCyanines have appeared upon the market under the names Éthyl Red, Homocol, Isocol, Orthochrome T, Pericol, Pinachrome, Pinachrome Violet, Pinaverdol and Sensitol Green, whilst cyanines were marketed under the names Ethyl Cyanine T and Quinoline Blue.

The first suggestion as to the constitution of cyanines and isocyanines was made by Decker, who had discovered the alkylquinolinium pseudobases and regarded these as intermediates (Bor. 1891, 24, 690). Miethe and Book formulated the isocyanine molecule as having two quinolino nuclci linked in the 2:4'-positions to a :CH group (Ber. 1904, 37, 2008), but with two extra hydrogen atoms as suggested by Spalteholz. It was W. König who pointed out how unlikely is their presence and he gave | 1 See p. 530 for explanation of alternatiformulæ for cyanine (I) and isocyanine (II) clature for this and subsequent compounds.

$$\begin{array}{c} \begin{pmatrix} 7 & 6 \\ 8 & 5 \end{pmatrix} & \begin{pmatrix} 6' & 7' \\ 5' & 8' \end{pmatrix} \\ RN1 & 4 \end{pmatrix} = \begin{array}{c} 9 \\ C H - \begin{pmatrix} 4' & 1'NR1 \\ 3' & 2' \end{pmatrix} \end{array}$$

1:1'-Dialkylcyanine iodide. Bis-(1-alkyl-4-quinoline)methincyanine iodide.1 I.

1:1'-Dialkylisocyanine iodide. (1-Alkyl-2-quinoline)(1-alkyl-4-quinoline)methincyanine iodide.

TT.

(J. pr. Chem. 1906, [ii], 73, 100). König made the alternative suggestion that in cyanine and isocyanine one quinoline ring may have undergone fission to an open chain (ibid. 1912, [ii], 86, 166), but this was disproved by Mills and Evans's synthesis of o-aminocinnamylidenequinaldine methiodide, in structure very close to the open-chain formula for isocyanine, but differing in properties (J.C.S. 1920, 117, 1035), and further evidence against it was advanced by König and Treichel (J. pr. Chem. 1921, [ii], 102, 63). Vongerichten and Höfchen produced evidence that the 2-methyl group of quinaldine is concerned in isocyanine formation (Ber. 1908, 41, 3054), whilst Kaufmann and Vonderwahl proved that the 4-position of the quinoline nucleus is involved (Ber. 1912, 45, 1404). The constitution of isocyanine (III) was definitely established by Mills and Wishart, by its oxidation to 1-alkyl-2-quinolone and cinchonic acid alkochloride (J.C.S. 1920, 117, 579); whence that

1:1'-Dlalkylisocyanine iodide. (1-Alkyl-2-quinoline)(1-alkyl-4-quinoline)methin-cyanine iodide.

III.

of cyanine follows. The formula of isocyanine is confirmed by Adams and Haller's preparation of various isocyanines by the action of alkali on various lepidine alkiodides (J. Amer. Chem. Soc. 1920, 42, 2389), and by Hamer's synthesis from lepidine ethiodide and 2-iodoquinoline ethiodide (J.C.S. 1928, 206). Sho subsequently prepared complex isocyanines from 5-methylacridine methiodide and 2-iodo-

¹ See p. 530 for explanation of alternative nomen-

quinoline alkiodule, but these were not sensi-tieres (I C.S. 1930, 995). For accounts of the 1913. By effecting the condensation under preparation of secoganines, by the action of modified condutions, Hamer prepared a number caustic alkal on alcoholus solutions of quaternary of 22-cyanines and noted that these also are salts of substituted quinolines and quinaldines, and of their properties, including optical and photographic properties, see Sheppard (JCS. 1909, 95, 15), Wise, Adams, Stewart and Lund (Ind. Eng. Chem. 1919, 11, 460), Mills and Pope (Phot. J. 1920, 60, 183), Barbier (Bull Soc. chim. 1920, [iv], 27, 427), Hamer (JCS 1921, 119, 1432; Phot. J. 1922, 62, 8), and Sakuran (Bull Inst. Phys. Chem. Res. Japan, 1932, 11, 681, 1044); Hamer prepared complex tsocyanines from quinoline alkiodide and 2 methyl acenaphthpyridine methiodide (JCS

In naming isocyanines, plain numerals refer to the outpaldine nucleus, whilst those of the quinoline nucleus are followed by a dash 1933). (Mills and Pope, Phot J. 1920, 60, 183) Mills and Wishart's formula for teocyanane (III) differs from Konig's (11) in that the acid radical is attached to the alternative nitrogen atom, with a corresponding difference in the conjugated chain uniting the nitrogen atoms Mills and Wishart auggested that the two formulæ virtually tautomerio compounds represent (J.C.S. 1920, 117, 579), and virtual tautomerism has since been shown to be a characteristic of cyanines of various types (Mills and Braunof cyannes of various type of this and braun-holtz, thd. 1922, 121, 1459, Hamer, total 1928, 206, Kuhn, Winterstein and Balser, Ber 1930, 63, BB, 3176, Ogata, Proc Imp. Acad. Tokyo, 1932, 8, 119, Bull Inst. Phys Chem. Rcs. Japan, 1934, 13, 549; Brooker and Keyes, J. Amer Chem. Soc 1935, 57, 2488) Although the acid radical in cyanines is not now regarded as attached to one nitrogen atom, to the exclusion of the other, static formula are given throughout this article, for the cake of clearness. For the convenience of naming definite examples, the amon in the formula is usually represented as iodide, although other anions are equally applicable.

The cyanifies, to use the term in its general sense, are monacid salts which can conteniently be crystallised from methyl or ethyl alcohol, but are only slightly soluble in hot water. The erystalks usually contain solvent of crystalks a recent publication denies this (Lauer and tion; they have a metallic lustre and are often [Horio, J. pr. Chem. 1935, [1], 143, 303). The pleochrone. Although the solutions are de-formula (V) for pinacyanol was suggested by

aalts are unstable.

Although the 44's and 24'-eyammes have been known from such an early date, the pre-paration of 2.2'-eyanine (IV) by Fischer and

1·1· Dimethyl ψ cyanine fodide Bis (1·methyl 2 quinoline)methincyanine iodide

Scheibe, by condensing 2 iodoquinoline methcontacts of quandino methioduc (J. pr. Chem. and who found that on oxidation it green 1920, [19], 100, 80) as much more recent. Instead quandinic acid and 1-th): 2 quandons (J. C.S. of their term ½-isoquarine, ½-lills and Oams 1920, 117, 1550). Their formula was confirmed.

photographie sensitisers (J.C.S. 1928, 206). For the preparation of more complex & cyanines. 2-todo β naphthaquinoline alkiodide uss condensed with an alkiodide of quinaldine or β naphthaquinaldine (Hamer and Kelly, 1bid. 1931, 777), and 2 iodoquinoline alkiodide was condensed with 2-methylacenaphthpyridine methodide (Hamer, thid. 1930, 995). The I.G. Farbenind, A.G. prepared the diethyl analogue of IV from 1 ethyl 2 thioquinelone ethiodide and quinaldine ethiodide in pyridine (B.P. 423792, 1932), whilst Kendall prepared IV from 2-methylthiolquinoline, methyl n. toluenesulphonate, and malonic acid in pyrkline. followed by potassium iodide (B.P. 431141.

22'-, 24', and 44'.Carbocyanines -The most valuable sensitiser of its day, conferring photographic sensitivity even to red light, was tho hlue dye Pinacyanol, which was discovered by Homolka and patented by the Farbwerke vorm. Moister, Lucius and Bruning (G P. 172118, 1905) (later known also as Quinaldina Blue and Sensitol Red., Naphthacyanole, Pinachrome Blue and Sensitol Violet belong to the sama class) It was prepared by the action of caustio alkali on an alcoholic solution of quaternary salts of quinoline and quinaldine, in the presence of formaldehyde. The part played by formaldobyde may also be taken by a trihalogenomethene (G P. 200207, 1907) or by chloral hydrato (Ogata and Suzuki, Bull. Inst. Phys Chem Ros Japan, 1934, 13, 488) The preparation of pinseyanol and its homologues (" & dicyanines"), by the formaldehyde method, is described by O. Fischer (J. pr. Chem 1918, [11], 88, 204), Wise, Adams, Stewart and Lund (Ind. Eng. Chem. 1919, 11, 480), Mills and Pope (Phot. J. 1920, 69, 253), Braunholtz (J.C.S. 1922, 121, 169), Moudgill (totd. 1922, 121, 1509), Mers and Gutekunst (Ind. Eng Chem. 1922, 14, 1060), and Hamer (J CS. 1923, 123, 2333). Inscher observed that the quinoline salt takes no direct part in the condensation (l c.); although it has always been accredited with increasing the yield, pleochroic. Although the solutions are deformula (V) for pinacyanol was suggested by colorised by excess of acid, the colourless diacid. Mills and Pope (I c) and established by Mills and

Bis (1-ethyl-2-quinoline)trimethincyanine todide.

Hamer, who concluded from analysis that the compound is formed according to the equation 2C10H2N C2H2X+CH2O

 $=C_{25}H_{25}N_2X+H_2O+HX+2H$

by König's preparation from quinaldine ethiodide and cthyl orthoformate in the presence of acetic anhydride (Ber. 1922, 55, [B], 3293); Hamer, by using pyridine instead of acetic anhydride, obtained 70-79% yields by this method (J.C.S. 1927, 2796), and further con-firmed formula V by converting methylenediquinaldine dialkiodide (VI) into (V), by the

Methylenediquinaldine dialkiodide.

action of alkali in the presence of a quinolinium salt (J.C.S. 1923, 123, 246), and by reducing (V) to (VI) (J.C.S. 1925, 127, 211); a process for converting (VI) into (V) by treatment with an oxidising agent in a non-alkaline medium was subsequently patented (Soc. Chem. Ind. in Basle, B.P. 321155, 1928). Rosenhauer with Hoffmann and Unger (Ber. 1926, 59, [B], 946) obtained the methylene bases from quinaldinium salts in the solid state. By condensing such a base with iodoform, the 2:2'-carbocyanine (V) was synthesised by Fischer and Rosenhauer (Z. angew. Chem. 1923, 36, 330) and by Rosenhauer, Schmidt and Unger (Ber. 1926, 59, [B], 2356). Kuhn and Winterstein prepared 2:2'-carbocyanines, otherwise difficult to obtain, by the action of acetic anhydride and sodium formate on the appropriate methylene base, or on its salt with an organic acid (B.P. 328357, 1929). Sakurai has described the sensitising action of certain 2:2'- (and 2:4'-) carbocyanines (Bull. Inst. Phys. Chem. Res. Japan, 1932, 11, 1044), and Kendall the preparation of a 2:2'-carbocyanine, from 2-methylthiolquinoline, p-toluenesulphonate, glutaconic acid, and pyridine (B.P. 431141, 1933). Ogata, Kawasaki and Masuda prepared a 2:2'-carbocyanine containing two nuclei isomeric with phenanthroline from a salt of the appropriate base, ethyl orthoformate, and acetic anhydride (Bull. Inst. Phys. Chem. Res. Japan, 1934, 13, 486); for preparing a simple 2:2'-carbocyanine, Ogata and Suzuki treated quinaldine methiodide with chloral hydrate and alkali (ibid. 488).

When the constitution of pinacyanol was established as (V), Mills and Hamer predicted that, corresponding with this 2:2'-carbocyanine, there would also be a 2:4'- and a 4:4'-carbocyanine (VII and VIII) (J.C.S. 1920, 117, 1550) This prediction was soon fulfilled, for Mills Braunholtz synthesised (VIII) (J.C.S.

1:1'-Dialkyl-2:4'-carbocyanine salt. (1-Alkyl-2-quinoline)(1-alkyl-4-quinoline)trimethincyanine salt. VII.

1:1'-Dialkyl-4:4'-carbocyanine salt. Bis-(1-alkyl-4-quinoline)trimethincyanine salt. VIII.

whilst Mills and Odams 1923, 123, 2804), 1913). synthesised (VII) (ibid. 1924, 125, Thus Mills and Braunholtz showed that when quinaldine ethonitrate is condensed with di-oformylalkylaminodiphenyl disulphide (IX), in pyridine solution, one product is (V), the disulphide having furnished the central carbon

Dl-o-formylalkylaminodiphenyl disulphide.

atom of the three-carbon chain; when lepidine ethonitrate is used instead of quinaldine ethonitrate, the corresponding product must be They identified this dye with Krypto-(VIII). cyanine (also called Rubrocyanine), which Adams and Haller had prepared by using a lepidinium instead of a quinaldinium salt in the carbocyanine condensation (J. Amer. Chem. Soc. 1920, 42, 2661), and which has proved a valuable sensitiser for infra-red light. By interaction of lepidinium salts, pyridine and ethyl orthoformate, Hamer prepared 4:4'-carbocyanines (J.C.S. 1927, 2796), together with neocyanine (q.v.). Mills and Odams, by the action of (IX) on a mixture of the ethonitrates of quinaldine and lepidine, with a continuous the second of the continuous three continuous transfer of the con synthesised the 2:4' carbocyanine (VII). From a spectroscopic examination they concluded that the dicyanine, which had been prepared from quaternary salts of 2:4-dimethylquinoline and its derivatives (G.P. 155541, 1903; Fischer, J. pr. Chem. 1918, [ii], 98, 204; Wise and Adams, Ind. Eng. Chem. 1918, 10, 801; Mikeska, Haller and Adams, J. Amer. Chem. Soc. 1920, 42, 2392; Palkin, Ind. Eng. Chem. 1923, 15, 379), is a 2:4'-carbocyanine.

Thiacarbocyanines, Thiacyanines, Thia-2'cyanines (Thia-\psi-cyanines), and Thia-4'-cyanines (Thiaisocyanines).—Hofmann observed that a purple dye, which he regarded as the cyanine of the benzthiazole series, is formed by heating an aqueous solution of the alkiodides of benzthiazole and 2-methylbenzthiazole with ammonia (Ber. 1887, 20, 2251). Mills showed that the reaction proceeds more readily in pyridine and isolated a yellow compound in addition to the purple one. Analyses led him to the surprising conclusion that the yellow dye is the cyanine (X), and the purple dye the carbocyanine (XI), of the benzthiazole series, and this was confirmed by synthesising the thiacyanine (X). (XI) sensitised a gelatino-bromide and (X) a gelatino-chloride photographic emulsion (J.C.S. 1922, 121, 455). The name thia-

Bis (3 ethyl 2 benzthiarole)methincyanina rodide.

3 3'-Diethylthiaenrhocyanine fodide Bis (3 ethyl-2-benzthiazole)frimethincyanine rodide

cyanine is in accordance with international nomenicature and therefore preferred to the original term "thiocyanine" Similarly, the original name "carbathiocyanine" for (XI) was first replaced by "throcarbocyanine" (Hanier, J C.S 1927, 2796), and has now become thin carbocyanine Tho thiacyanino (X) corresponds with ψ eyamine, isocyamine, and eyamine, and the thurcarbocyamine (XI) with the 22-, 24-, and 4 4'-earbocyanines. Just as the disulphido (IA) was used to synthesise 22'-, 24', and 4.4'carbocyanines, so, similarly, by condensing it with 2 methylbenzthiazolo ethiodice, Mills and Braunholtz obtained (XI); they lound that, in addition to the function of (IX) in providing the central carbon atom of the three carbon chain of a carbocyamine, it also furnishes a benz thiazolo nucleus, which condenseo with the quaternary sait available to give a dyo in which the nuclei are linked by one CH- group; thus from (IX) and 2-methylbenzthiazole ethiodide, the other product is (X) (ibid. 1923, 123, 2804) Fisher and Hamer prepared thiacyanines from

a 2 methylkenztharolum or a 1 methyl-anaphtathaxolum chlordo, by the action of amyl mitrite in the presence of acetic anhydrido, G.U.S. 1930, 2502). The I of. Farhennd, A. C. applied this method to the preparation of thiacyanines with complex nuclei (B P. 4000). 1931, and also described aminothiacyanines (B P. 4000). 1931, Monig. Klest and Gétzo prepared thaxyanines, including one (XII) of conuderable complexity, by treatment of 3-nethylbenzthiarolone, or its derivatives, with magnesium mitthji lodde (Ber 1931, 64, fB).

3 4 3',4'- Bis-trimethylenelhlacyanine lodide Bis (3 4 trimethylene-2 benitharole)methinepanine (odide. XII.

1604). Kendall prepared thingyannes by bent, \$\frac{\text{Hancoles (B I \cdot \text{2.7887}, 1922), and of 2-menty)}{\text{Beta}} and \$\frac{\text{Hancoles (B I \cdot \text{2.7887}, 1922), and of 2-menty)}{\text{Beta}} and \$\text{Beta}\$ beta prepared than \$\text{Constant}\$ \text{Constant}\$ by the accuracy of \$\text{Beta}\$ and \$\text{Log}\$ beta prepared than \$\text{Constant}\$ \text{Constant}\$ by the accuracy of \$\text{Beta}\$ and \$\text{Log}\$ minutes by the accuracy of \$\text{Beta}\$ and \$\text{Log}\$ minutes by the accuracy of \$\text{Log}\$ and \$\text{Log}\$ minutes by the accuracy of \$\text{Log}\$ and \$\text{Log}\$ minutes by the accuracy of \$\text{Log}\$ and \$\text{Log}\$ minutes by \$\text{L

azolono alkiodule with 2 methylbenuthazolo alkiodule in pyridime (B.P. 427302, 1923). Closely related is Kendall's method, in which a quaternary sait of 2-methylbenuthazolo; no cents with one of 2 methylbenuthazolo; no claims that it is advantageous to make the preparation and condensation of the saits all one process (B P. 427509, 1933).

" Thruzole Thracarbocyanines (X1 Purple") were prepared by Konig and Menr by condensing a 2-methylbenzthazolium sult with ethyl orthoformate, or anhydrous sodium formate, in the presence of various solvents (J. pr Chem. 1925, [u], 109, 324) Hamer found the ethyl orthoformate and pyridine method to be most successful in the case of the thiacarboeyanmes (J C.S 1927, 2796), and subsequently con densed ethyl orthoformate, in the presence of pyridine, with salts of methyl-o- and β naphtha thiazoles (d C.S. 1929, 2598), as also did Brooker and Kodak Ltd. (B 1', 351261, 350163, 1029). The I G Parbenned, A. G. used ethyl orthoformate and pyridine in preparing thincorbe cyanines with complex nuclei (B.P. 300217, 1931) and in preparing certain substituted this carbocyanines (B 1' 418745, 420071, 418745. thiacarbocyanines (B 1) 418745, 420071, 1932; 421015, 1933), including those from methyltetrahydro a and Bunjihthathiazolis (BP 427887, 1932) and from 2 methyl \$\beta\$-naphtbathmzole (B.P 452409, 1934) | Kendali prepared thisearbocypnines by heating together methyltholbenzthiazole, alko.p.toluene sulphonate, crotonio anhydride, or glutacome acid, and pyridine (B P. 431141, 431186, 1933).

Mdls and Braunholtz found that when the dasalphado (1x) reacts with quinalism ethonitate, it behaves as oxplained under this eyanines, so that the 22'-carbo(yanino (V) is accompanied by the thin2' cyanine (XIII) (I C S. 1923, 123, 2804) Hamer confirmed

3 t - Dialkylthia-2'-cynnine iodide (1-Allyl 2 quanolene)() allyl 2 benthazole)meU in eyanine todide XIII.

this formula by mepsing this 2-yannes by the action of caushie alkan on the alkudides of 2-sodopunohine and 2 methylterathraziole (the 1928, "200), whist more complex this-2-yannes were prepared by mang methyle and napathathrazole alkudides and 2 sodo-6-naphthacquinohine alkudide (thet. 1920, "2551 Hamer and Kib), thet. 1931, 737 The 1G. Farbenind. A. G. subsequently condensed 2-sodogunohine alkudides with saits of still more complex thinacle derivatives (B.P. 400161, 1931), of methylterahydro-and \$\theta\$ purplicathrazoles (B.P. 427887, 1932), and 62-methyl 1931), of methylterahydro-and \$\theta\$ purplicathrazoles (B.P. 427887, 1932), and 62-methyl printine on a mixture of 1-ctiv)-2 thou action of printine on a mixture of 1-ctiv)-2 thou municipal alkudides and 2 methylberathrazoles alkudides and 3 methylberathrazoles and 3 methylberathrazol

dall found it better to use alkyl-p-toluenesulphonates of the thiono compounds (B.P. 438420, 1934); he also prepared thia-2'cyanines by condensing salts of I-methylthiolbenzthiazole and quinaldine (B.P. 424559, 1933). Beilenson and Hamer observed that introduction of chlorine or bromine atoms into the thia-, thiacarbo-, or thia-2'-cyanine molecule has little effect upon the absorption maximum (ICCS 1998 1995)

(J.C.S. 1936, 1225).

Mills and Braunholtz found that when the disulphide (IX) reacts with lepidine ethonitrate, the 4:4'-crarbocyanine (VIII) is accompanied by thia-4'-cyanine (XIV) (J.C.S. 1923, 123, 2804), which they had previously obtained by the

3:1'-Dialkylthia-4'-cyanine iodide. (1-Alkyl-4-quinoline)(3-alkyl-2-benzthiazole)methincyanine iodide. XIV.

action of caustic alkali on the alkiodides of 2-methylbenzthiazole and quinoline (ibid. 1922, **121**, 2004). Brooker and Kodak Ltd. condensed quinoline alkiodide with alkiodides of methyl- α - and β -naphthathiazoles (B.P. 378455; 380140, 1930), whilst the I.-G. Farbenind. A.-G. used salts of still more complex thiazole derivatives (B.P. 400401, 1931), of methyltetrahydro-a- and β -naphthathiazoles (B.P. 427887, 1932), and of 2-methyl-ββ'-naphthathiazole (B.P. 452408, 1934). The I.G. Farbenind. A.-G. prepared thia-4'-cyanines hy the action of pyridine and piperidine on alkiodides of lepidine and 3-methylthiohenzthiazolone (B.P. 423792, 1932), and Kendall prepared them hy condensing quaternary salts of lepidine and 2methylthiolbenzthiazole, or, conversely, of 4methylthiolquinoline and 2-methylbenzthiazole (B.P. 424559, 1933).

Fuchs pointed out that certain sensitisers containing henzthiazole nuclei are superior to the older cyanines in that they inhibit chemical fog (Chem.-Ztg. 1933, 57, 853); this is also confirmed by Sobolev, Bondareva and Evteeva, who compared the sensitising action of various substituted cyanines containing benz- and naphtha-thiazolo nuclei (J. Appl. Chem. Russia, 1936, 9, 335).

OTHER CYANINES AND SYMMETRICAL CARBOCYANINES.

Mills and Smith's discussion of the reactivity of the methyl group in heterocyclic bases (J.C.S. 1922, 121, 2724) is of especial interest in connection with those bases which have since been found to give eyanine dyes. In the present section, the plan has been adopted of dealing, in roughly chronological order, with each base which was employed, on account of its reactive methyl group, for eyanino dye formation; this

dall found it better to use alkyl-p-toluene- is followed by an account of further types made sulphonates of the thiono compounds (B.P. possible by methods depending on the reactivity 438420. 1934): he also prepared thia-2'- of groups other than methyl.

Indocyanines, Indocarbocyanines, and Indo-2'-cyanines.—W. König pointed out that an indolenine salt such as (XV), can condense

2:3:3-Trimethylindolenine methiodide. XV.

methyleneindoline.
XVI.

with ethyl orthoformate, in the presence of acetic anhydride, to give the indocarbocyanine (XVII) ("Astraphloxine") (Ber. 1924, 57, 685), which method was subsequently used by Ghigi (Gazzetta, 1933, 63, 698). Its sensitising properties were noted by Hamer, who prepared

1:3:3:1':3':3'-Hexamethylindocarbocyanine iodide. Bis-(1:3:3-trimethyl-2-indolenine)trimethincyanine iodide.

XVII.

indocarhocyanines by use of ethyl orthoformate and pyridine (J.C.S. 1927, 2796). They may also be prepared by condensation of the methylene hase (XVI), which corresponds with (XV), with formaldehyde (Soc. Chem. Ind. in Basle, B.P. 312174, 1928). Kuhn and Winterstein prepared them by interaction of acetic anhydride, sodium formate, and a methylene hase, or its salt with an organic acid (B.P. 328357, 1929). Ogata prepared (XVII) hy treatment of (XV) with chloral hydrate and alkali (Proc. Imp. Acad. Tokyo, 1933, 9, 602). Wahl and the I.G. Farbenind. A.-G. made "Indolenine Yellow" by treating salts such as (XV), or the corresponding hase (XVI), with an ester or salt of nitrous acid, in the presence of acetic anhydride (G.P. 459616, 1926). Kuhn, Winterstein and Balser pointed out that neither the analytical results nor the colour of the compound accorded with the formula proposed, and they established that it is in fact the indocyanine (XVIII). By treating an acetic acid solution of

1:3:3:1':3':3'-Hexamethylindocyanine Perchlorate. Bis-(1:3:3-trimethyl-2-indolenine)methineyanine perchlorate.

XVIII.

roughly chronological order, with each base (XVI) with sodium nitrite and then with perwhich was employed, on account of its reactive incthyl group, for cyanino dye formation; this (XIX), and showed that this reacts with

3 3-Dimethyl 2 aldoxime indolenine methoperchlorate
XIX.

(XVI), in acctic anhydride solution, to give that undocyanne (XVIII); an unenpumetrotal radio-cyanne was also prepared (Ber. 1930, 63, [B], 310). The intermediate (XIX) was also obtained by the action of natroxyl chlorids on (XV) or (XVI) (I.G. Farbennd, A.G., BF, 309733, 1939). Nitroxylubphane and may be used similarly (I.C. I. Ltd., Lanch, and Piggott, BP, 391215, 1931) Indo 2° cyanines (XX)

1 3 3 1' Tetramethylindo 2' cynnine iodide. (1 3 3 2 rimethyl 2 indolenine/1-methyl 2-guinoline)methincydnins iodide

were prepared by reaction of (XV) with 2 odo quinohne alkinodie in the presence of casette alkalı (Hamer, J.C.S. 1928, 206), but tretchyiagme it recommended as a better consideraagme (Kockk, Ltd., B. 2 408371, 1832). With general consideration of the second of the sacid, the midocyanines and indicarbocyanines give colour lakes which are fast to light, etc. (L.G. Farbenind, A. G., B. P. 448508, 1934)

Ozacyanines, Ozacarbocyanines, cyanines, and Oza 4' cyanines - König end Meier were the first to use 2 methylbenzoxazole methiodide for the preparation of a cyanine, condensing it with ethyl orthoformate and condensing it with early to a constraint and a control analysis of the control of using ethyl orthoformate in conjunction with pyridine and noted that the oxacarbocyanines have photographic sensitising properties (J.C.S. 1927, 2796). Oxacarbocyanines were similarly prepared from methyl a and β naphthoxazoles by Brooker and the Eastman Kodak Co. (U.S.P. 1939201, 1933; 1962123, 1934) and from 2 methyl $\beta\beta'$ uaphthoxazola by the I.G. Farbennd, A.G. (B.P. 432969, 1933). Kendall prepared oxacarbocyamnea by reaction of the 2 methylthiolbenzonazole alko p-toluene-sulphonates with glutacome acid (B P. 431141, 1933), or with crotonic anhydride, and pyridine (B.P. 431186, 1933). The colourless exacyanines were prepared by Fisher and Hamer, by the action of amyl nitrite and acetic anhydride on aslts of 2 methylbenzoxazole, and their pale yellow analogues were similarly prepared from methyl a. and B.naphthoxazoles (J.CS. 1934, Kendall prepared ovacyanines by tha action of malonic acid on a 2-methylthiolbenzoxazole alko p-toluenesulphonato in pyridina (B.P. 431141, 1933), and was able to prepare both symmetrical and unsymmetrical axacyanines by reaction of a 2 methylbenzovazole quaternary salt with one of 2-methylthiolbenzorazole

(B P. 424559, 1933) or of 3-alkyl-2 thiobenzovasolone (B.P. 438420, 1934). The I.G. Farben. ind. A.-G. obtained oxa-2 cyanines by condens-ing quaternary salts of 2 methylbenzoxazole and 1-alkyl 2-thioquinolone (B.P. 423792, 1932; B.P. 423827, 1933) and patented oxa 2'- and oxa 4'-cyanines containing naphthoxazole nuclei (B.P. 388898, 1931; B.P. 432969, 1933). Kendall prepared oxa-2'-cyanines from salts of 2 methylthiolbenzovazole and quinaldine (BP. 424559, 1933). Brooker and Keyes found that, whereas the yield of oxa-2' cyanine obtainable by the ection of caustic alkali on an alcoholic solution of the ethiodides of 2-methylbenzoxazola and 2 iodoquinoline is very low, it can be greatly increased by using a strong organic base es condensing agent (J. Amer. Chem. Soc. 1935, 57, 2488). The parent oxa, oxacarbo. oxa-2'-, and oxa 4'-cyanines are represented by formulæ (X), (XI), (XIII), and (XIV) respec-tively, if each sulphur etom be replaced by oxygen.

Belenacganines, Belenacarbocyanines, Selena-2'-cyanines, and Selena-4'-cyanines .- Clark first used 2 methylbenzaelenazole quaternary solts for the preparation of cyaninea, and by their condensation with ethyl orthoformate and pyridme he prepared sclenacarbocyanines, which he found to be sensitisers (JCS 1928, 2313; of I.G. Farbenind. A.G., B.P. 383486, 1931; 418745, 1932). From such salts or their derivatives, the I.G. Farbenind. A.G. prepared, hy the action of amyl nitrite and acetic anhydride, selenacyanines (B P. 380702 400951, 1931), by condensation with quinolinium aalts, sciena-4'-cyanines (B.P. 389969, 400951, 1931; cf. U.S.P. 2051134, 1936), and by condensation with a salt of 2 lodoquinoline or 1-slkyl-2 thioquinolone, siena 2. cyanines (B P. 386791, 400951, 1931; 403840, 423702, 1932). The parent selena selena-carbo, selena-2. and selena-4 cyanines may he represented by formula (X), (XI), (XIII), and (XIV) respectively, if each sulphur atom be replaced by eclenium.

2 1. Cyanina, 4 1. Cyaninas, This 1. Cyaninas, and Sekan. 1. Cyaninas, — This and Hame used 1. adoisequanoline alkindude, in place of 2. adoingouseline alkindude, in place of 2. adoingouseline alkindude, for cyanine condensations. Thus with quandinium salts in the presence of alcohol and alkal it gave 2. 1. Cyaninas (XXI), which were also obtained in much lower yield, by condensation of the alkindudes of inoquinione and quinaldine. They possed out that isoQuinoline Red, which had been prepared by interaction of isoquinoline, quandline, benotirelicitied, and since olthoride

1 2 Dialkyl 2.1 cyanics fodide.
(1-Allyl 2-quinoline) (2-allyl 1-isoquinoline) methincyanine todide

XXI.

isoQuinoline Red. 1:2'-Benzylidene-2:1'-cyanine chloride. 1:2'Benzylidene(2-quinoline)(1'-isoquinoline)methincyanine chloride.

XX11.

isoquinoline alkiodide was further condensed with lepidine alkiodide to give a 4:1'-cyanine (XXIII); with salts of 2-methylbenzthiazole,

$$\begin{array}{c|c}
\hline
 & 7 & 0 \\
8 & 5 \\
\hline
 & 8' & 5 \\$$

1:2'-Dialkyl-4:1'-cyanine iodide. (1-Alkyl-4-quinoline)(2-alkyl-1-isoquinoline)-methincyanine iodide.

XXIII.

etc., to give thia-1'-cyanines; and with 2-methylbenzselenazole alkiodide to give a selena-l'cyanine. Whereas the absorption maximum of each l'-cyanine closely approximates to that of the analogous 2'-cyanine, the photographic sensitising action of the former is much weaker than that of the latter (J.C.S. 1934, 1905). Brooker and Keyes, by taking advantage of the reactive methyl group of salts of 9-methylphenanthridinc, which may be considered as a derivative of both quinoline and isoquinoline, prepared four cyanines. In each instance the absorption maximum lay nearer the blue end of the spectrum than that of the isomeride containing a β -naphthaquinoline nucleus, whilst no sensitising action was shown (J. Amer. Chem. Soc. 1936, 58, 659).

Thiazolocyanines, Thiazolocarbocyanines, Thiazolo-2'-cyanines, Thiazolo-1'-cyanines, Thiazolo-4'-cyanines, and Thiathiazolo-cyanines. -Mills and Smith pointed out the reactivity of the methyl group in a 2-methylthiazolium salt and, by condensing 4-phenyl-2-methylthiazole methiodide with quinoling methiodide, ohtained the 4-phenyl derivative of the parent thiazolo-4'cyanine (XXIV) (J.C.S. 1922, 121, 2724; cf.

3:1'-Dialkylthiazolo-4'-cyanino iodide. (3-Alkyl-2-thiazole)(1-alkyl-4-quinoline)methincyanine iodide.

XXIV.

carbocyanine (XXV) together with the corre-

3:3'-Dimethylthiazolocarbocyanine iodide Bis-(3-methyl-2-thiazole) trimethincyanine iodide.

XXV.

sponding thiathiazolocyanine (J.C.S. 1923, 123, 2804). Fisher and Hamer prepared thiazolo-2:4-dimethylthiazolium carhocvanines from salts by the pyridine and ethyl orthoformate method (J.C.S. 1930, 2502). Brooker and Kodak Ltd. showed that, as prepared by the latter method, these are accompanied by the corresponding neocyanines (q.v.) (B.P. 408273, 1931). Kendall prepared a thiazolocarhocyanine from 2-methylthiolthiazolium p-toluenesulphonate and malonic acid in pyridine (B.P. 431141, 1933). Thiazolo-2'-cyanin's (XXVI), prepared

3:1'-Dialkylthiazolo-2'-eyanine iodide. (3-Alkyl-2-thiazole)(1-alkyl-2-quinoline)methin-cyanine iodide.

XXVI.

from 2-methylthiazolium and 2-iodoquinolinium salts, were described by Brooker, Keyes, and White (J. Amer. Chem. Soc. 1935, 57, 2492; cf. Kodak Ltd., B.P. 408569; 408570, 1932; I.G. Farbenind. A.-G., B.P. 386903, 1931), and thiazolo-1'-cyanines, similarly prepared from 1-iodoisoquinolinium salts, were described by Fisher and Hamer (J.C.S. 1934, 1905).

Oxazolo- and Selenazolo-2'-cyanines.—By condensing 2-methyl-oxazolium or -selenazolium salts with 2-iodoquinolinium salts, Brooker, Keyes and White prepared oxazolo-2'-cyanines and a selenazolo-2'-cyanine (J. Amer. Chem. Soc. 1935, 57, 2492). They may he represented hy formula (XXVI), if the sulphur atom he replaced hy oxygen and selenium respectively.

Thiazolin ocarbocyanines, Thiazolino-2'cyanines, and Thiazolino-4'-cyanines.—Quaternary salts of 2-methylthiazoline were first applied to cyanine formation hy Brooker, who hy condensation with ethyl orthoformate and pyridine ohtained thiazolinocarbocyanines, with 2-iodoquinolinium salts ohtained thiazolino-2'cyanines, and with a quinolinium salt obtained a thiazolino-4'-cyanine, all of which are photo-graphic sensitisers (J. Amer. Chem. Soc. 1936, 58, 662). The formulæ of these three types resemble (XXV), (XXVI), and (XXIV) respec-tively, but the thiazoline nucleus in each instance takes the place of the thiazole nucleus. As compared with the corresponding dyes containing the thiazole nucleus, those with the thiazoline nucleus have their absorption maxima nearer to the region of short wave-length

(Brooker, I c.). Selenazolinocarbocyanines.-White and Kodak Ltd. similarly used quaternary salts of 2methylselenazoline to get selenazolinocarbo-

cyanines (B.P. 392410; 406074, 1931). The formula resembles (XXV), but the selen azolino nucleus takes the place of the thrazole

nucleus

2 2' and 2 4' Pyridocarbocyanines, other Pyridocyanines, and Miscellaneous Cyanines .- Pyrido carbocyanines cannot be obtained from pico linium salts by the general methods used for other carbocyanines, but Rosenhauer and Barlet succeeded in preparing 2 2 pyridocarbocyanines (XXVII) by treating 2 picoline quaternary salts, in alcoholic solution, with potassinm hydroxide and chloroform; by using a a picolinium salt, they prepared the 4-4 pyrido-carbocyanine (XXVIII) (Ber. 1929, 62, 2724) Ogata and Suzuki prepared (XXVII) by

1 1' Dialkal 2 2' pyridotarboczanine iodide Bis (1-alkyl 2 pyridine)trimethineyanine rodide XXVII.

treating an a picolinium salt with chloral bydrate and alkalı (Bull Inst. Phys Chem. Res. Japan, 1934, 13, 488) Hamer and Kelly were able to get 2 pyrido 2'-cyanines (XXIX)

1 I'- Dialkyi 2 pyrido 2' cyanine lodide (1 Aliyi 2 pyridine) I-aliyi 2 quindine prethineyanıne todide.

by the action of caustic alkali on alkiodides of applied to the preparation of indothia-, indoza, 2 iodopyridine and quinaldine, but not by the oxograpio, oxiding, and oxidinazolo cyanines alternative method of using those of a picoline (B.P. 424559, 1933). By condensing alkyl pand 2 iodoquinohue (J.C.S. 1931, 777). Brooker tolurnesulphonates of 3 methylthiobenzoxazoand Keyes, by employing thethylamme as condensing accent, however, were successful the can 2 "pyridacyenine (XXXII) by the second method, and O for S), and this method of using ally in improving its yield by the first method, p toluenesulphonates of a thione and of a and found this reagent superior to caustic alkah methyl- substituted base was also applied to for 2' cyanine preparations in general, and were some of the types mentioned above (BY. thus enabled to make cyanines not previously 438420, 1934). accessible, such as 2 pyrido 4'cyanine, 4 pyrido.
2' cyanine (XXX), 2 2' pyridocyanines (XXXI), than those containing a quinoline or isoquinoline 435542, 1933). Brooker, Keyes and White It is convenient to arrange the prefixes in

1 1'-Methyl 4 pyrido-2'-cyanine perchlorate (1-Ethyl 4 pyridine)(1-ethyl-2-quinoline)methin cyanine perchlorate, XXX.

1 1'-Diethyl 2 2'-pyridocyanine collide Bu (1 ethyl 2 pyridine)methineyanine codide, XXXI.

used triethylamine in condensing 2 iodonyridinsum saits with 2-methyl thiazohum or selenazolium salts, to give 2 pyridothiazolocyanines (XXXII) and 2 pyridoselenazolocyanine (XXXII, substituting Se for S) (J. Amer. Chem. Soc.

3 1' Diethyl 2'-pyridothiazolocyanine iodide. (1 Ethyl 2 pyridine)(3 ethyl 2 thiazole)methin-cpanine iodide XXXII.

1935, 57, 2492), whilst 2 'pyridothiazolino cyanines are similarly obtainable (Brooker and Kodak Ltd , B P. 437807, 1933). By con densing quaternary salts of a picoline and 2-methylthiolbenzthiazole, Kendali prepared 2-pyridothiocyanine (XXXIII) and this general method of applying a reactive methylthiol

3 1'-Dialkal 2' pyridothiaevanine lodide (t-Allyl 2 pyridine)(3-nilyl 2 beneihiazole)methinchanine logids. XXXIII.

in conjunction with a methyl group was also

and 24-pyridocyanines (J. Amer. Chem. Soc. nucleus, confusion is avoided by using plain 1935, 57, 2488). Bellenson and Kodak Ltd. numerals to refer to the nucleus which is menfound potassium carbonate to be superior to tioned first, whilst the numerals with a dash caustic alkalı for eyanıne condensations (BP. refer to that which is mentioned second.

oxa- precedes thia -.)

Of other nuclei which have been introduced into cyanines, Hamer, Heilhron, Reade and Walls first used the quinazoline nucleus, preparing a quinazocarbocyanine from a 2-methyl salt, ethyl orthoformate and pyridine (J.C.S. 1932, 251). Kendall, hy taking advantage of reactive alkylthiol and methyl groups, prepared various types of cyanines containing one or two heterocyclic nuclei having two nitrogen atoms, such as quinazoline, pyrimidine and thiodiazole (B.P. 425609, 1933). By condensing the alkylthiol derivative of such an alko-p-toluenesulphonate with malonie acid in pyridine, he obtained symmetrical cyanines, in which two nuclei of this kind are linked by :CH. (B.P. 431141, 1933). With erotonic anhydride in pyridine a carhocyanine containing two thiodiazole nuclei was similarly prepared (B.P. 431186, 1933). Ogata described a carhocyanine containing two henziminazole nuclei as heing prepared from 1;2-dimethylhenziminatory. azole methiodide, diphenylformamidine, potassium acetate, and acetic anhydride (Proc. Imp. Acad. Tokyo, 1933, 9, 602).

Bases, of which Cyanines are the Quaternary Salts .- Mills synthesised the ethiodide of dibenzthiazolylmethane, which, by elimination of acid, gave the base (XXXIV), of which

 ${\it 3'-} Ethylbenzthiazolenyl-2-benzthiazolylmethane.\\$ XXXIV.

(X) is the ethiodide (J.C.S. 1922, 121, 455). 2-imino-3-methylbenzthiazoline from and quinaldine, prepared the base, of which thia-2'-eyanine (XIII) is the alkiodide (J.C.S. 1936, 507). Kendall patented a series of such bases, including those of which 2:2'-, oxa-2'-, selena-2'-, thia-2'- and thia-4'-eyanines also indo-2 earhoeyanine, are the quaternary salts, the method being to condense a hase, having a reactive methyl group, with a quaternary salt, having a reactive alkylthiol, or substituted aminovinge group; the resultant bases sensitise differently from, and sometimes more powerfully than, the corresponding salts (B.P. 456362, 1935).

Unsymmetrical Carbocyanines.—The methods which have been described so far for preparing earbocyanines lead of necessity to products in which the two heteroeyelie nuclei are identical. It is true that mixtures of two quaternary salts may he used, as in the preparation of 2:4'-earbocyanines, but in this case the desired product must be separated from the isomerides which accompany it. Apart from this, Mills and Raper were the first to prepare unsymmetrical 2:2'-carhoeyanines, by heating the methylene hase from quinaldine ethiodide with a substituted quinaldinium salt and formaldehyde in alcoholie solution (J.C.S. 1925, 127, 2466). The Society of Chemical Industry in Basle claimed that by condensing 1 mol. of a compound X-CH:NH (e.g. formimino ethyl ether hydrochloride) with 2 mols. of a heterocyclic ammonium salt con-

alphabetical order, e.g. indo- precedes oxa-, and taining a reactive methyl group, or with the corresponding pseudo-hase, symmetrical carhoeyanines result, whilst equimolecular quantities give an intermediate, which may subsequently be condensed with a second mol. of methylene hase to give an unsymmetrical carhocyanine. Examples are the intermediate prepared from (XVI), also those from certain quinaldinium salts; the first of these may he condensed with a second mol. of (XVI) to give (XVII), or with a different methylene base to give an unsymmetrical carhocyanine (B.P. 334706, 1928). I.C.I. Ltd., Piggott, and Rodd similarly used diarylformamidines, ArNH-CH:NAr, to get either symmetrical carhocyanines (when 2 mols. of quaternary salt containing a reactive methyl group are treated with one of

ArNH-CH:NAr),

or intermediates (when equimolecular proportions react). Thus (XV) heated with diphenyl-formamidino gives (XXXV), whilst in the presence of acetic anhydride its acetyl derivative (XXXVI) is produced (B.P. 344409, 1929).

2-ω-Anilinovinyl-3:3-dimethylindolenine methiodide. XXXV.

 $2-\omega$ -Acetanilidovinyl-3:3-dimethylindolenine methiodide. XXXVI.

They found that such intermediates, by heating with acetic anhydride, potassium acetate, and a second mol. of heterocyclic ammonium salt containing a reactive methyl group, give earbo-cyanines, for example, the intermediate from diphenylformamidine and quinaldine ethiodide gives with (XVI) the unsymmetrical indo-2'carbocyanine (XXXVII); that derived from 2-methylhenzthiazole ethiodide gives with (XV) the indothiacarbocyanine (XXXVIII); (XXXVI) with 2-methylhenzoxazole ethiodide gives the indoxacarbocyanine (XXXIX); and the intermediate derived from 2-methylhenzthiazole ethiodide gives with 2-methylhenzoxazole ethiodide the oxathiacarbocyanine (XL); these four dyes are photographic sensitisers (B.P.

1:3:3-Trimethyl-1'-ethylindo-2'-carbocyanine lodide. (1:3:3-Trimethyl-2-indolenine)(1-ethyl-2-quinoline)-trimethincyanine iodide. XXXVII.

1 3 3 Trimethyl 3*-ethylindothlacarbozzanine iodide (1 3 3*Trimethyl-2-indoleune)(3-ethyl 2 benzthuzole)trimethincypnine todide.

XXXVIII.

1 3 3-Trimethyl 3'-ethyllndoxacarboe; anine

(133 Trimethyl 2-indolenine)(3 ethyl 2 benzoxazole) trimethincyanine iodide XXXIX.

3 3*Dieth) loxathiar processine iodide (3 Ethyl-2 benzozatel)(3 ethyl 2-bensthearde)from thingyanine rodide XI.

354899, 1930) Ogata applied their method to preparation of various carbocyanines demonstrated the virtual tautomerism of this aeries; whereas the dye prepared from a quaternary salt A and the intermediate from a salt B is identical with that prepared from the salt B and the intermediate from A, the yields may be different. He obtained better results with the acetanihdo than with the anihnointermediate. By use of dimethylquinoline, etc, he prepared carbocyanines with an extra methyl group, which was subsequently caused to react with p dimethylaminobenzaldehyde (Proc. Imp. Acad. Tokyo, 1932, 8, 119; Bull Inst. Phys. Chem. Res Japan, 1934, 13, 501, 519, 549, 556). In preparing an indoquinazocarboevenine, Kendali used pyridine, in place of acetic anhydride and potassium acetate (B.P. 425609, 1933). The IG Farbenind. A.G states that indoxacarbocyanines are especially good sen-sitisers (B.P. 442160, 1933) To prepare carbo cyanines containing an indolenine nucleus, they condensed an aldehyde, such as (XLI;

1 3 3-Trimeth; lindoline 2 methylene-waldeb; de XLI.

(R.P. 438278, 1933), with a quaternary salt congroup. 9.Ellys! and 9 phenyl-steinoarthocyanine taning a reactive methyl group (or with the and 9 style concritosyanine were also decrete corresponding methylme base), in the presence (J. Amer. Chem. Soc. 1935, 57, 2480). White of a solvent and condensing agent (B.P. 438603, the condensing agent (B.P. 438603, the condensity of the condensity

Gyunines with Substituents on the Chain-The first recorded instance of a cyanine substatuted on the chain is the duntro-derivative which Mills and Hamer obtained by treating 2.2-carbocyanine with nitric acid (J C S. 1930, 117, 1550; Hamer, tid. 1928, 3160), and that accord is necoganize, which will be dealt with in a later section. A direct synthesis of the 10 phenyl 2.2-carbocyanine (XLII) was

10-Phenyl 1 1-dimethyl 2 2' carbocy anine chloride. Bis (1 methyl 2 guinoline) phenyltrimethineyanine chloride.

XLII.

effected by Fascher and Rosenhauer (Z angew. Clem. 1923, 36, 303), and by Nosenhauer, Schmidt and Unger (Ber. 1928, 59, 2356), by heating the methylene base from quantidam methomethyl-asiphate with hencetrichloride in alcohole soleton (ef. Ogast and Soutis, Bull Inst. Phys. Chem. Res. Japan, 1934, 13, 479). Hinner, by methyl group with ethyl orthoacette in the presence of pyridine, succeeded in preparing a nethyline described processing the succeeded in preparing a nethyline described processing and the succeeded in preparing a nethyline described processing and the succeeded in preparing a nethyline described processing and the succeeded in the presence of pyridine, succeeded in preparing a nethyline described processing and the succeeding and the su

Nethyl 3 3' dialkylthiacarbocy anine iodide
 Bis (3-all pl 2-benzthiazole)β methyltrimethiacyanine
 vodide.

XLIII.

being negative with other types of heterocyclic bases (J.CS. 1928, 3160). Subsequently 9methylthracarbocyanines were obtained from 1-methyl a- but not from 2-methyl & naphthathiszohum iodides (ibid 1929, 2598). Brooker and White, however, obtained a 9 methylthracarboeyanine from 2 methyl 8 naphthathis role, by using the etho-p toluenesulphonate instead of the ethiodide of the base. They prepared 9 methylselenscarbocyanine and 9. methylazacarbocyanine by use of a 2 methylbenzoxazolium modide, ethyl orthoacetate, and trethylamine in pyridine solution. observed that 9-methylthiacarbocyanine is formed on heating a 2-methylbenzthinzolum salt with pyridine only, and that this same method is applicable to salts of methyl aand 8-naphthathiazoles, 2 methylbenzselenazole, and 2-methylbenzoxazole (J. Amer. Chem. Soc. 1935, 57, 547). By condensing 1-methyl-benzthiazohum salts in the presence of pyridine with various ortho-esters, they prepared thiscarbocyanines in which the central carbon atom of the chain carried an alkyl, aryl, or aralkyl group. 9-Eihyl- and 9 phenyl-selenacarbocyanine

prepared 7-ethylthiazolinocarbocyanines (J. Amer. 1 the rings improved either the photographic Chem. Soc. 1936, 58, 662). Arranging the thiacarbocyanines in order of increasing wavelength of absorption maximum, Brooker and White found the 9-substituents to fall in the order Me, Et, H, Ph, and the 4:5:'4:5'- and 6:7:6':7'-dibenzthiacarbocyanines and selenacarbocyanines were similar, but the oxacarbocyanines abnormal (J. Amer. Chem. Soc. 1935, 57, 2480); the same order was later found for 6:6'-dicbloro- or -dibromo- substituted thiacarbocyanines (Beilenson and Hamer, J.C.S. 1936, 1225). The I.G. Farbenind. A.-G. has patented meso-substituted oxa-, selena-, thia-carbocyanines, which are variants of those described above, through having either more complex nuclei or nuclei carrying substituents (B.P. 396217; 402458; 410481, 1931; 415949, 420971; 421015; 418745; 427887, 1932; 432969, 1933; 452408, 1934). Although they have patented dyes prepared by the action of carbon tetralialogenide on a methylcne base derived from a methyl-indoleninium salt (e.g. (XVI), derived from (XV)), or from the corresponding 2-methylbenzthiazolium or quinaldinium salts, no formula is given (B.P. 373160, 1931). Kendall and Ilford Ltd. patented the preparation of meso-substituted carbocyanines by using an imino-ether, R'O·CR:NH, instead of an ortho-ester (B.P. 404997, 1932); also by condensing a quaternary salt, containing a reactive methyl group, with an acid anhydride, in the presence of an acid-binding agent (B.P. 369236, 1930).

Ogata found the 9-methyl group of thiacarbocyanines to be reactive: by treatment of a 9-methylthiacarbocyanine with furfuraldehyde, benzaldehyde, or p-dimethylaminobenzaldehyde in the presence of alcohol and piperidine, he prepared the corresponding 9-furylvinyl-, 9-styryl-, 9-p-dimethylaminostyrylthiacarbocyanine (XLIV) (Bull. Chem. Soc. Japan, 1936, 11, 262).

3:3'-Diethyl-9-p-dimethylaminostyrylthiacarbocyanine iodide.

Bis-(3-ethyl-2-benzthiazole) \$\beta-p\-dimethylaminostyryltrimethincyanine iodide. XLIV.

By making use of the ortho-ester of thiophena-carboxylic acid, the I.G. Farbenind. A.-G. prepared carbocyanines with an a-thienyl group in the meso-position (B.P. 403845, 1932). They also introduced, into this position, groups such as cyclohexyl-, furyl-, or pentadecyl-, by treating a benzoxazolium salt, containing the desired group in the 2-position, with sodium ethoxide in alcohol, followed by a quaternary salt containing a reactive methyl group (B.P. 439359, 1933). By means of ortho-esters, RS-CH2-C(OEt)3, the RS·CH₂· group was introduced into the 9-position of thiacarbocyanines, but neither this nor the introduction of alkylthiol groups into

sensitising or the stabilising action of the parent dyes (Kiprianov, Suitnikov and Suitsch, J. Gen. Chem. Russ. 1936, 6, 576).

The foregoing methods lead to meso-substituted symmetrical carbocyanines, but the I.G. Farbenind. A.-G., by utilising the alkyl ester of an arylated thioimide, ArN:CR·SR', prepared unsymmetrical carbocyanines with the substituent R in the meso-position (B.P. 412309, 1932).

König, Kleist and Götze, by condensing 2-ethylbenzthiazole methiodide with orthoformate, isolated a thiacarbocvanine (XLV), in which the two lateral carbon atoms of the chain bear substituents (Ber. 1931, 64, 1664).

3:8.10:3'-Tetramethylthiacarbocyanine salt Bis-(3-methyl-2-benzthiazole)ay-dimethyltrimethin-cyanıne salt.

XLV.

The I.G. Farbenind. A.-G. applied such quaternary salts, in which the reactive methyl group is replaced by -CH₂R, to the preparation of other symmetrical and unsymmetrical carbocyanines with one or more R groups in the a-position in the chain (B.P. 405309; 411876, 1931), and also to obtaining 2'-cyanines, e.g. XLVI, in which the nuclei are linked by :CMe. (B.P. 405309, 1931;

8:6'-Dimethyl-3:1'-diethylthia-2'-cyanine iodide. (6-Methyl-1-ethyl-2-quinoline)(3-ethyl-2-benzthrazole)methylmethineyanine iodide.

XLVI.

Götze, however, reported 423792, 1932). that when ethiodides of 2-ethylbenzthiazole and 2-iodoquinoline are treated with alkali, the 2'-cyanine is accompanied by an unknown dye (Angew. Chem. 1936, 49, 563).

By heating the alko-p-toluenesulphonate of a base containing an alkylthiol group (or, alternatively, the isomeric thione) with cyclopentadiene and pyridine, Kendall prepared carbocyanines in which the three-carbon chain, joining the heterocyclic nuclei, itself forms part of a cyclopentadiene nucleus. The method was applied to the preparation of carbocyanines from dimethylthiolthio- $\beta\beta'$ -diazole, and of 2:2'-carbocyanines (e.g. XLVII), as well as of oxa- and thia-

1:1'-Dimethyl-9:11-vinylene-2:2'-carbocyanine iodide. Bis-(1-methyl-2-quinoline)ay-vinylenetrimethin-cyanine iodide.

XLVII,

carbocyanines. By using indene instead of | Piggott, and Rodd isolated an intermediate cyclopentadiene, an o phenylene group may be compound, which, by condensation with a introduced instead of vinylene, and this has been heterocyclic ammonium salt containing a

2 2° Dimethylthiol-1 1° diethyl 5 7 o phenylene-6 6° pyrimidacarbocyanine bromide Bis (2 methylthiol 1° ethyl 6 pyrimidine)ay-a phenylene-trimethineyanine bromide XLVIII.

the nuclei, the compounds sensitise, or even desensitise, chloride or bromide emulsions, the cyclisation of the chain has a hypsochromic effect (B.P 431142; 431187, 1933).

Dicarbocyanines - Besttie, Heilbron Irving condensed a heterocyclic ammonium salt containing a reactive methyl group (or the corresponding methylene base) with derivatives of hrome, chlore or mitro malendialdehyde, or a suitable acraldehyde derivative, ϵg the anil of a bromo β anilmoacraldehyde, PhN CH CBr CH NHPh, either in pyridine in the presence of piperidine, or in acetic anhydride In the presence of potassium accetate; they thus prepared oyanines in which the two heterocyclic nuclei are linked by a five carbon chain, the central carbon atom of which carries as substituent Br, Cl, or NO. As eyanines with a three-carbon chain had been designated " carbo cyanines," these new dyes were called "dicarbo cyanines" By using alkindides of quinaldine, Jepidine, 2 methylbenzthiazole, 233 trimethylin dolerane, and 2 methylbenzoxazole, there were ohtained, respectively, meso substituted 2 2' . 4 4'. thia., indo., and oxa dicarbocyanines (e g XLIX)'

Bis (1 3 3 trimethyl 2 indolenine)y nitropentamethin-cyanine iodide.

XLIX.

They found that whilst the halogenodicarbocyanines are sensitisers for the deep red dmitrophenylpyridmium chloride (LI), or (b) the or near infra red, the corresponding nitrodi or near mine rest, see Corresponding mixed by Enth Obsailes, (by Enth Obsailes and Wilson acrobocyanines possess describting properties Annalen, 1907, 338, 107) by treating (LI) with a (J.C.S. 1932, 260; cf. I.C.I. Ltd., Heilbron, and Irring, B. P. 3,5388), 1939). By the action of & snilmoacraldchy de anil,

PANH CH CH CH.NPh.

in acetic anhydride, on an equimolecular quantity of a salt such as (XV), I.C.I. Lt.1,

done with ora, thus, 22°, 22°, pyrido, quanza-, reactive methyl group, gave a dioarboyama and 4.4 pyrindo carboyamae (e.g. XIVIII), with an unsubstituted five carbon chan, besides others containing two this \$\beta^{\beta}\]. or this-. Where the two quaternary salts are the same, \$\beta^{\beta}\]-disazion bucklet. According to the nature of the resultent dye is symmetrical, where they differ it is unsymmetrical; eg the inter-mediate from (XV) with 1-methylbenzthiazole ethiodide gave (L), or, alternatively, this was

(1 3 3 Trimethyl 2 indolenine)(3 ethyl 2 benzthiazole)pentamethineyanine rodide,

prepared from the intermediate obtained from 2 methylbenzthiazole ethiodide, together with (XVI) (BP. 555693, 1930). For preparing dicarbocyanino intermediates, the I G. Farbenand, A G, found at advantageous to use an organio base, in alcohol, as condensing agent; they recorded the absorption maxima of many such intermediates, and the sensitising and absorption maxima of various types of un-symmetrical dicarbocyanines prepared from them (BP 434234; 434235, 1933) They have patented various substituted dicarbocyanines, including dyes carrying one or more mothyl groups on the chain (B.P. 388204; 394837, 1931; 427887, 1932, 452408, 1934) and others are described by Bellenson and Hamer (J CE 1936, 1225). Kendall and Hford Ltd. used propargylaldehyde acetal, or its derivatives, for condensing with quaternary salts containing reactive methyl groups, to furnish the three central carbon atoms of the dicarbocyanise chain (B.P. 300508, 1931), and prepared a thiadicarbocyanine by heating 2 methylthiol henzibiazole metho p toluenesulphonato with sorbic anhydride in the presence of pyridine (BP, 431186, 1933). In Ogata's preparation of dicarbocyamnes by heating a carbocyamne intermediate, e.g. (XXXV), and a quaternary salt (XV), with orthoformic ester and acetic anhydride, the course of the reaction is not clear (Proc. Imp Acad. Tokyo, 1934, 10, 572).

Tricarbocyanines .- Tricarbocyanines, in which the nucles are linked by a seven carbon chain, were described by three sets of workers. Wahl and the I G. Farbenind, A .G. condensed (XV) or (XVI) with (c) a compound such as 24 dye (LII) obtained (by Zincke and Würker,

2 4 Dinitrophens lpyridinium chioride.

5-Methylanilino-1-phenyliminomethochloride- $\Delta^{2.4}$ pentadiene.

base such as methylaniline. With equimolecular proportions, in acetic anhydride, (LIII) resulted. But 2 mols. of quaternary salt to 1 mol. of (LI) or (LII) gave the indotricarbocyanine (LIV) (G.P.

MePhN·CH:CH·CH:CH·CH:CH·RI

6-Methylanilino-△1:3:5-hexatriene-3';3'-dimethy-lindolenyl methiodide.

LIII.

1:3:3:1':3':3'-Hexamethylindotricarboeyaninc iodidc. Bis-(1:3:3-trimethyl-2-indolenine)heptamethincyanine iodide.

LIV.

499967, 1928). I.C.I. Ltd., Piggott, and Rodd also condensed (XVI) with a compound of type (LII) to give (LIV), which they observed to be a sensitiser for infra-red light (B.P. 355693, 1930). Fisher and Hamer prepared dycs by use of (LI) or a compound of a type similar to (LII), preferably the latter, carrying out the reaction in alcoholic solution in the presence of caustic alkali; the generality of the method, in that various quaternary ammonium salts containing a reactive methyl group are applicable, was pointed out, and the sensitising properties for infra-red light were noted. There were prepared thiatricarbocyanines (LV), including those containing α- and β-naphthathiazole nuclei, selenatricarbocyanines (as LV but Se for S), thiazolinotricarbocyanines, and 2:2'-tricarbocyanines (LVI). The indotricarbo-

R1:CH-CH:CH-CH:CH-CH:CH-R2

$$R^{1}$$
 is $C < ; R^{2}$ is NRI

2:2'-Dialkyithiatricarbocyanine iodide. Bis-(2-alkyl-1-benzthiazole)heptamethincyanine vodide.

R1:CH-CH:CH-CH:CH-CH:R2 R¹ is ; R² is NR NRI

1:1'-Dialkyi-2:2'-tricarbocyanine iodide. Bis-(1-alkyl-2-quinoline)heptamethincyanine iodide. LVI.

cyanine (LIV) was obtained by use of acetie anhydride and sodium acetate (J.C.S. 1933, 189).

amine, etc., instead of caustic alkali, for con-densing an intermediate of type (LII) with the appropriate quaternary salt, succeeded in preparing thiazolotricarbocyanines and 4:4'tricarbocyanines (Xenocyanine) (LVII) (B.P.

R¹·CH·CH:CH·CH:CH·CH:CH·R² ; R² is -VRÍ R¹is RŃ

1:1'-Dialkyl-4:4'-tricarbocyanine iodide. Bis-(1-alkyl-4-quinoline)heptamethincyanine iodide. LVII.

436941; 437017, 1933). Brooker, Hamer and Mees published spectrograms illustrating the sensitising action of tricarbocyanines (Phot. J. 1933, 73, 258); as did also Dieterle, Dürr and Zeh, who pointed out that various types of di-and tri-carbocyanines had been independently prepared in Germany, chiefly owing to the work of König, and put into commercial use (Z. wiss. Phot. 1933, 32, 145). Certain more complex, or substituted, tricarbocyanines have been described (I.G. Farbenind. A.-G., B.P. 388204, 1931; 452408, 1934; Beilenson and Hamer, J.C.S. 1936, 1225). As a tricarbocyanine intermediate, Kendall found 2-benzoxazolylpyridinium chloride, prepared from 2-chlorobenzoxazole and pyridine, to be preferable to (LI) (B.P. 424264, 1933). Brooker and Kodak Ltd. patented the preparation of tricarbocyanines by interaction of a quaternary heterocyclic ammonium salt, containing a reactive methyl group, with an alkiodide of 2-iodo-pyridine or -quinoline, in pyridine solution (B.P. 435252, 1933).

By making use of an intermediate, similar to (LII), but prepared from a suitably substituted pyridinium salt, the I.G. Farbenind. A.-G. synthesised tricarbocyanines carrying a methyl on the chain (B.P. 394537, 1931). Corbellini and Fusco halogenated anils of type (LII) and used these to make tricarbocyanines with a halogen atom on the chain, and found these dyes to be as good sensitisers as the parent dyes (Rend. Ist. Lomb. Sci. Lett. 1935, [ii], 68, 961). Dieterle and Zeh record, on the other hand, that tricarbocyanines carrying a p-acetoxy group on the chain (I.G. Farbenind. A.-G., B.P. 441624, 1933) (see also under tetraand penta-carbocyanines) are less good sensitisers than the unsubstituted compounds (Z. wiss. Phot. 1935, 34, 245). To prepare tricarbocyanine intermediates, similar to (LIII), the I.G. Farbenind. A.G. found an alkaline medium advantageous; they recorded the absorption maxima of several such intermediates, and the sensitising action of various unsymmetrical tricarbocyanines prepared from them (B.P. 438449; 438450; 438484, 1933).

Comparisons of Series .- In the indo- series, Kuhn and Winterstein found that the tricarboeyanine is the most easily reduced, the dicarbocyanine less readily, and the carbocyanine still less readily, the cyanine being the most resistant Brooker and Kodak Ltd., by using triethyl- (Ber. 1932, 65, [B], 1737). In the 2:2'-, 4:4'- indo-, oxa-, pyrido-, thia-, thiazolino- and thia-zolo- series. Ogata tabulated the melting point of the corresponding carbo, dicarbo, and triearbo-cyanine, and the censitising maximum of each ; ho found the latter to be shifted regularly towards the red on lengthening the polymethin chain. Ho also compared the yields when corresponding carbo, dicarho- and tricarbo eyanines are prepared by use of the appropriate and in an acid or alkalino medium respectively (Proc. Imp. Acad. Tokyo, 1932, 8, 421; 1933, 9, 602,
 Buil, Inst. Phys. Chem. Res. Japan, 1934, 13, 528;
 Ogata and Kimura, ibid. 1934, 13, 537). Tho optical and photographic properties of typical cyanines were studied by Bloch and Hamer (Phot. J. 1928, 68, 21; 1930, 70, 374) Fisher and Hamer compared the frequencies of maximum absorption of methyl alcoholic solutions of the cyanine, carbe-, dicarbe- and tracarbo-cyanino of various symmetrical types, anhydride or chloride in the presence of pyridine, correlating frequency changes with changes in the stability is, however, increased: the prechemical constitution; eg the values of the paration of compounds such as (LA) (where n= frequency-decreases, caused by each successive [0, 1, 2 or 3) made possible the preparation of lengthening of the chain joining the nuclei, became smaller with increasing length of chain (Proc. Roy. Soc. 1936, A, 154, 703)

Tetra- and Penta carbocyanines—Somewhat

as the pyridinum nucleus of (LI) had been ruptured to give the open-chain compound (LII), so König had ruptured the furfural nucleus, by treatment with an aromatic base, to give (eg) (LVIII) (J. pr Chem. 1905, (u), 72, 655) and that of furylacrolem to give (e g.) (LIX) (n=1) (ibid, 1913, (u), 88, 193)

[PhNH CH CH CH C(OH)CH NPh]HBr

a Hydroxyglutaconaldehyds diantiide hydrobromide

LVIII.

Subsequently, König (with Hoy, Schulze, Silberkweit and Trautmann) prepared correaponding dyes with nine- and eleven membered carbon chains (L1X, n=2 and 3, respectively), by rupture of the nuclei of 5-furyl \(\Delta^{24}\) penta-dieno-1-al and 7-furyl \(\Delta^{246}\) heptatrieno 1-al These dyes are unstable, the respectively. instability increasing as the chain is lengthened, By acylation (or benzoylation), with on acid

R CH CH CH C(OAc) (CH.CH),-CH.R1

cyanines in which the nuclei are joined by longer chains (Ber. 1934, 67, [B], 1274) The I.C. Farbenind. A.G. has patented such cyanines, with a chain of at least seven carbon atoms, and describes them as excellent agustisers to infrared biht: the examples include 19 acetoxythia-

10-Acetoxy-6 6'-diphenyl 3 3'-dimethylthiatricarbocyanine perchlorate Bus (6-phenyl 3 methyl-2 bensthuszole) y scelozyheplamethincyanine perchlorale. LXL

13-Acctoxy-1 1'-dimethyl 4 4'-tetracarbocyanias perchlorate
Bis (1-methyl-4-quinoline)e-acetoxymonamethincyanins perchlorate LXII.

13-Acetoxy-1-1'-dimethyl 2 2'-pentacarbocyanine perchiorate.
Bu-(1-methyl 2-quinoline)e-acetoxyundecamethineyanine perchiorate.

corresponding acetoxy- indo-, selena-, and 4:4'tetracarbocyanines (e.g. LXII), also 11-acetoxythia- and 13-acetoxy-2:2'-pentacarhocyanines

(e.g. LXIII) (B.P. 441624, 1933).

Brooker and Keyes have published spectrograms showing the sensitising action of such tetra- and pentacarhocyanines, hy means of which the furthest photographic excursions into the infra-red have heen made (J. Franklin Inst. 1935, 219, 255). For descriptions of the sensitising action of dyes of this type, having chains of various lengths, see also Dieterle and Zeh (Z. wiss. Phot. 1935, 34, 245).

More recently Dieterle and Riester have succeeded in synthesising tetra- and pentacarhocyanines not substituted in the chain; these are more powerful sensitisers than their acetoxy-substituted derivatives (Z. wiss. Phot.

1937, 36, 68, 141).

apoCyanines.-By heating quinoline alkiodide with caustic alkali in alcoholic solution, Kaufmann and Strühin ohtained as chief product a red compound, erythroapocyanine, together with some yellow compound, xanthoapocyanine; these were unlike the other cyanines then known, in heing more stable to acid (Ber. 1911, 44, 690; cf. G.P. 154448, 1903). König formulated them as (LXIV) and (LXV) respectively.

1:1'-Diethyl-3:4'-apocyanine iodide. (1-Ethyl-3-quinoline)(1-ethyl-4-quinoline)apocyanine iodide.

LXIV.

1:1'-Diethyl-3:2'-apocyanine iodide. (1-Ethyl-2-quinoline)(1-ethyl-3-quinoline)apocyanine iodide. LXV.

but without giving any evidence (Ber. 1922, 55, [B], 3293). The formulæ were, however, proved to he correct by Mills and Ordish, who, hy the action of potassium permanganate on the erythroapocyanine from quinoline ethiodide, obtained the same oxidation product as Kaufmann and Strühin, and, regarding the product as the diethiodide of a diquinolyl, they represented the reaction thus:

 $(C_{18}H_{13}N_2Et_2)I+I_2=(C_{18}H_{12}N_2Et_2)I_2+HI$

By analogy with other cyanines they concluded that the only two diquinolyls possible were those having the 3:4- (LXIV) and 3:2-linkings (LXV)

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tricarhocyanines (e.g. LXI), 12-acetoxy- (or respectively, thus supporting König's formulæ. 12-benzoxy-) thiatetracarhocyanines and the Their views were confirmed by the synthesis of 3:4'-diquinolyl and the demonstration that its diethiodide is identical with the product obtained hy oxidising erythroapocyanine with iodine (J.C.S. 1928, 81).

Neocyanines.-Clarke and the Eastman Kodak Co. observed that when lepidine ethiodide is treated in alcoholic solution with iodoform and alkali, there is formed, in addition to the 4:4'carhocyanine (VIII), a less soluble dye giving a green solution (U.S.P. 1804674, 1931). This compound, neocyanine (or "allocyanine"), sensitised further into the infra-red than any dye then known (Dundon, Schoen and Briggs, J. Opt. Soc. Amer. 1926, 12, 397; Bahcock, Nature, 1928, 121, 830). Hamer found that neocyanines also accompany (VIII) when lepidine quaternary salts are heated with ethyl orthoformate in the presence of pyridine; hy varying the conditions, the formation of one dye at the expense of the other could be favoured, and a 40% yield of neocyanine was thus obtained. It was shown analytically that the molecule contains two iodine atoms, three lepidine residues, and either one or two extra carbon atoms; on the hasis of one extra carbon atom, a formula was postulated (J.C.S. 1927, 2796; 1928, 1472). Ogata described similar dyes in which the three lepidine nuclei are replaced by quinaldine or a-picoline nuclei, and studied the effect of various reagents upon neocyanine formation (Proc. Imp. Acad. Tokyo, 1932, 8, 503; Bull. Inst. Phys. Chem. Rcs. Japan, 1934, 13, 491; Ogata and Tamura, ibid. 475). Of these reagents, succinic acid proved interesting, and he obtained a neocyanine, in 1% yield, by heating 2-methylbenzthiazole ethiodide with succinic acid and ethyl orthoformate, and also prepared a neocyanine from 1:2-dimethylbenziminazole methiodide (ibid. 1934, 13, 497). Meanwhile Brooker and Kodak Ltd. had observed that when a 2-methylthiazolium salt is heated with ethyl orthoformate, in the presence of pyridine, it behaves as does a lepidinium salt, in that the carbocyanine is accompanied by a dye of the neocyanine type, which is, in some instances, the main product; although the neocyanine is usually less soluble than the carhocyanine, the contrary is sometimes true. They postulated a formula for their neocyanine in which two acid radicals, three residues of methylthiazole, and two extra carhon atoms are involved (B.P. 408272, 1931; 408273, 1932). In König's formula for neocyanine, which may he simplified into (LXVI), the

Neocyanine ethiodide. LXVI.

number of extra carbon atoms is two; he which the thia azacyanine is the quaternary salt. regarded it as a substituted dicarbocyanine (Z. By condensing a heterocyclic nitrogenous base, wiss. Phot. 1935, 34, 15). From the absorption maxima of 44'-carbocyanine, 44' dicarbocyanine, and neocyanine. Hamer is of opinion attituded formamidines, eq. 2 aminobenethiazole that the last may be regarded more correctly as gave (LXX), which could be converted into the a substituted carbocyanine (Chem. and Ind. dazadisacerbocyanine (LXXI). Treatment of a 1935, 13, 640).

Azacyanines -Starting from 2-aminoquinobne and 2-chloroquinoline, Hamer synthesised the 2 2'-azacyanine (LXVII) (then called "22'azocyanine"), in which the linking between the two outnoine nuclei is not by .CH hat by :N .

1.1. Dimethyl 2 2 anseyanine lodide Bis (1 methyl 2 quinoline)azamethineyanine 10dide LXVII.

It differed from the cyanines in that it was not decolorised by acids (JCS 1924, 125, 1348). Fuchs and Grauaug synthesised carbooyanine analogues, in which the chain CH CH CH is replaced by .N N CH, by condensing a heterocyclic aldehyde (e.g. quinoline-2 aldehyde metho-perchlorate) with the hydrazone of a heterocyclio ketono (e g 3 ethyl 2 benzthiazolone hydrazone). They prepared two dyes containing one benzthiazole and one quinoline nucleus (LXVIII), one with two benzthiazole nuclei, and one with a pyridine and a benzthrazole nucleus

3 1'- Dialky! 8 9-diazathia 2 -carbocyanina perchlorate
(1-Alkyl 2-quinoline)(3-alkyl-2-benzihiazole)βybisasatrimethincyanine perchlorate. LXVIII.

They noted that the colours of these dyes in solution vary from yellow to deep red, and they all act as photographic desensitisers

(Ber. 1928, 61, [B], 57). Kendall, hy condensing a quaternary hetero-

cyclio ammonium salt, containing an NH, group in the a or y-position, with a similar salt, contoining a reactive alkylthiol group or iodine atom, prepared various 24'., 44'., ezs., thia. thia.2'., and thia-4'.azacyanines. When instead of a quaternary salt containing an NH, group, the corresponding base (e.g. 2 aminobenzthiazolo) was subjected to such a condensation (e.g. with a 2-methylthiulbenzthiazole quaternary salt), a hase, e g. (LXIX), was obtained, of

3'. Methyl 2 bensthiazolylideneaminobensthiazole, LXIX.

containing an a NH, group as substituent, with cthyl orthoformate, Kendall obtained sub-

2 2 Dibenzthia zolylformamidine. LXX.

2 3'-Dimethyl 8 10 diazathiacarbocyanine lodide But-(3 methyl 2 benzthrazole)ay-diazatrimethineyanine andide

LXXI. 2 ammobenzthiazolium salt with ethyl ortho-

acetate and pyridino gave the 9 methyl derivative of (LXXI) Kendall noted that (LXVII), (LXVIII), and their analogues, and also hases such as (LXIX) and (LXX), act as sonsitisers to gelatino chloride photographic emulsions (BF 447038, 447109, 1934; 456419, 1935) He has postulated the theory that a substituted amino group has the power of increasing the sensitising action of a cyanine dye when intro duced in such a position that the carbon chem between it and one of the original nitrogen atoms is odd as opposed to even; similarly the sensitising properties of (LXXI), as contrasted with the desensitising properties of (LXVIII), are attributed to replacement by an odd-numbered of an even-numbered carbon chain (Proc. Ninth Internat Congr. Phot, 1935, 227). Definition of a Cyanine -A cyanine is characterised by possessing two heterocyclic nucles containing nitrogen; one of these nstrogen atoms is tertiary and the other quaternary and they are united by a chain of con-jugated double bonds, so that the number of carbon atoms in the chain is necessarily uneven; one or more :CH- groups in the chain may be replaced by one or more nitrogen atoms.

Compounds related to the Cyanines -The above definition excludes certain types of compounds, which are sometimes described together with eyagines. Such are the following: (a) compounds in which both nitrogen atoms are quaternary and the chain consists of an even number of carbon atoms, (b) styryl compounds, to which type the sensitiser Pinaffavol belongs and to certain members of which type Konig (Z. wiss. Phot. 1935, 34, 15) assigned the name "semicyanines," (c) the related anils, which are usually desensitisers, and (d) dyes prepared by condensing certain cyanine intermediates with compounds containing a reactive ryclic or nonevelic methylene group.

Nomenclature of the Cyanines -According to the historical treatment adopted in the present article, each point of nomenclature has been

has grown up in the British and American literature has been made clear. As it requires acquaintance with such a large number of group names, it is, however, far from satisfactory; the cyanines are now revealed as comprising a far greater number of different types than was originally contemplated, and the nomenclature has also been complicated by the fact that some of the simplest members were not amongst the first to be prepared. König pointed out the inconveniences attaching to the use of these group names, and proposed a systematic nomenclature (Ber. 1922, 55, [B], 3293), which has, however, proved cumbersome in actual use. The system denoted in the 1936 edition of "Beilsteins Handbuch der organischen Chemie" (which deals with the literature up to 1910) has the great advantage that, once a few simple principles have been laid down, each and every cyanine can be named systematically. Such principles are as follows: (1) the above definition of a cyanine (which is narrower than that of Beilstein) is taken; (2) each ring is denoted by its usual description and numbering (following Richter's "Lexikon" and the "Journal of the Chemical Society"), and the positions of linking are indicated; (3) the number of methenyl groups linking the nuclei is denoted by apo (for none), methin, trimethin, pentamethin, etc.; (4) the presence of :N· replacing :CH· in the chain is denoted by the term aza, which is already in use in international nomenclature for :N· replacing :CH in a ring. Below tho graphic formula of each cyanine in the present article has been given the name at present accepted, whilst below that, printed in italics, has been added the systematic name as arrived at from the principles formulated above.

CYANOGEN v. CYANIDES (this vol., p. 503). CYANOMACLURIN, C₁₅H₁₂O₆. A colourless crystalline compound found in Jak-wood; it is fairly readily soluble in water and gives a colourless precipitate with basic lead acetate and a violet colour with ferric chloride. When fused with alkali it gives β-resorcylic acid and phloroglucinol. The penta-acetyl derivative, C₁₅H₇O₆Ac₅, has m.p. 136–138°. the pentabenzoyl derivative, m.p. 171–172°. Cyanomaclurin is probably a reduction product of morin (Perkin, J.C.S. 1905, 87, 715).

CYANOPHORIC GLYCOSIDES. A considerable number of glycosides have been isolated from plants which yield hydrogen cyanide when hydrolysed, whilst the formation of hydrogen cyanide by a plant is attributed to the presence of a glycoside. The most interesting of these is amygdalin (q.v.), which contains the sugar gentiobiose, as does also lotusin from Lotus arabicus. Vicianin, from the seeds of the wild vetch, contains the disaccharide vicianose.

dealt with as it arose, so that the system which as grown up in the British and American sin, sambunigrin, dhurrin, gynocardin, linamarin, literature has been made clear. As it requires hiptagin.

A list of plants containing hydrogen cyanide compiled by Greshoff (Rep. Brit. Assoc. 1906, 138) and another by Rosenthaler (Chem. Zentr. 1919, iii, 274; 1930, ii, 932) enumerate 360 varieties in 148 species and 41 families. In general the presence of hydrogen cyanide in plants appears to be incompatible with that of alkaloids and terpenes. The amount of oyanophoric glycoside varies according to the conditions. Linseed, which contains linamarin, produces considerably less hydrogen cyanide when grown in this country than in the Orient, where it is grown under conditions of drought and high temperature. The cyanophoric glycosides are hydrolysed by acids and by the specific enzymes which accompany them in the plants, most of them are attacked by emulsin.

CYANURIC ACID v. CYANIDES (this vol.,

GYCLAMIN. C₃₀H₁₀₀O₃₂. m.p. 251°, [a]_D -22°, the saponin glycoside present in cyclamen tubers (Cyclamen europaeum). According to Dafert (Ber. der Pharm. Ges. 1926, 264, 409; 1930, 268, 289), it is hydrolysed to the aglucone, cyclamiretin, G₃₅H₅₀O₅, glucose (3 mols.) and l-arabinose (2 mols.).

E. F. A.

apoCYCLENE.

apoCyclene, C_9H_{14} , m.p. 42·5-43°, b.p. 138-139°/764 mm., d_4^{40} 0·871, n_D^{40} 1·43144, is formed when camphenilyl chloride (I) is heated with diethylaniline or when camphenilol (II) is digested with potassium hydrogen sulphate at 180-190° (Komppa and Roschier, Annalen, 1922, 429, 187; Gratton and Simonsen, J.C.S.

$$H_2C$$
— CH — CMe_2
 CH_2
 H_2C — CH — CHR

I. R = CI ; II. R = OH .

 H_2C — CH — CMe_2
 CH_2
 H_2C — CH — $C:N·NH_2$

1935, 1621; Komppa and Nyman, Ber. 1936, 69, [B], 338). Since it is stable to potassium permanganate it can be readily purified. According to Nametkin and Alexandrov (Annalen, 1928, 467, 191), it can be conveniently prepared by heating camphenilone hydrazone (III) with mercuric oxide and alkali.

J. L. S.

CYCLINES (CYCLIC OLEFINS), The members of the different series are usually ribes substances do not for the most part over derived from eyche alcohols, which in turn are naturally, although the monocyche terpenes demonstrated as members of the group must be regarded as members of the group. They can, however, readily be derived from by various aimple derivatives of the saturation of the saturation of the group around the saturation of the saturation of the group could be allowed by the saturation of the group elbonds, subjuture scale, call of y passes of to be distinguished according to the number of carbon atoms forming the might of the saturation of the group elbonds, subjuture scale, call of y passes of the saturation of the group elbonds, subjuture scale, call of y passes of the saturation of the group elbonds, subjuture scale, call of the passes of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the group at the saturation of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the group elbonds, subjuture scale, call passes of the group elbonds, subjuture scale, call passes of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the group elbonds, subjuture scale, call passes of the saturation of the sat

Mono-olefins.—The number of known monoolefine; octens a very large, and the various the corresponding chlorides or bramide by creatment with haldes of hydrogen, phosphorus senes represented by these compounds ranges from that in which the ring contains four earth of a very large of their very large atoms (cyclobutenes) to that in which it contains between terms of the contains four earth of the contains the contains seventeen carbon atoms (cycloberptaderens). commonly by the passage of their vapour

Letones. The alcohols are dehydrated discriptly warming them or refluving them with small or larger amounts of suitable dehydration-catalysts (hydrobromic said, jodine, zine-chloride, sulphuric acid, etc.) or by passage of their vapours over suitable contact catalysts (alumina, kaolin, etc.) heated to appropriate into the corresponding chlorides or broundes by treatment with haldess of hydrogen, phosphorus not obelieve statement with haldess of hydrogen, phosphorus or sulphur, and thence nint the cycle olefin by the action of boding alcoholic canatic skila, boding quindless or broundes by commonly by the passage of flour apour the passage of their apour the passage of their apour and the passage of their apour in quantity by catalytic hydrogenation of the corresponding phenols (Sabatter Senderens process), and hence cyclobracenes in mentioned relatively chapted on commercial scale. In the case of certain, disklyd queforecame in mentioned the partial chapters are included the case of certain, disklyd queforecames in mentioned the type of the partial chapters and the sumpersion of the consequence of the partial chapters of the case of certain, disklyd queforecames (mentioned) the hydrogenation of the class of certain disklyd queforecame in mentioned) the hydrogenation of colonia terpenes (menthadicines).

of petroleum like odour and in their physical and chemical properties closely resemble the open chain olefins. Certain physical properties

Mono ollying Ciclenes

Вр (℃) d	FF p
cycloButene 15-2	0 733°	
,, Pentene , 41·1-44	6 0 77434 1	421818
,, Hexene , 83-83 :	5 0 810220	4451221
" Heptene 113-11:	5 0 822829 1	455220
" Octene 145	0 85520	·47425
" Pentadecene , mp 36-	37 -	
" Heptadecene mp 4	7] —]	
1-Methyl-∆1-cyclopentene 72	0 775820	_
1-Methyl △2- ,,		1-422214
1.1-Dimethyl-△² ,,		1-419020
1·2· ,, ·Δ· ,, · 103–103	3 5 0 792320	1-447135
1-Methyl △¹ cyclohexeno	0 809920	1 4496 ²⁰
1· ,, · \(\(\alpha^2 \), , , 104-10	6 -	_
1. ,, .∆³. ,, 103	0 798600	1 44316
1 2 Dimethyl-∆3 cyclohexene	65 0 822620	1 4580218
1:4- ,, -Δ1- ,,	0 7985224	1 444*2
1.1	12 0 8056162	1-445102
1·3· " ·Δ³- " · · · · · 124-12		1 446621
13	0 8074184	1-450517

of the more important eyclenes are given in the table (p. 532). The most important of the simple cyclenes are cyclopentene and cyclohexene, which are respectively the parent compounds of a large array of alkyl- and polyalkyl-cyclopentenes and -cyclobexenes. The alkyl- and polyalkylderivatives of both series are almost invariably obtained from the corresponding alcohols or their balogen esters by the methods mentioned above. Allied to the cyclenes and resembling them in methods of formation and in ehemical behaviour arc the alkylidene-cyclanes of general formula, $(CH_2)_n > C = CRR'$.

cyclo Propene is reported to be formed together with other products by heating barium pyromucate, and cyclobutene by the distillation of trimetbyl cyclobutyl-ammonium iodide, but the identity of both products appears to be some-

wbat uncertain.

cyclo Hexene is a typical cyclene and is prepared on a large scale by dehydration of cyclohexanol. When the latter (400 g.) is heated with concentrated sulpburic acid (12 c.c.) at 130-140°, and finally at 150°, cyclobexene distils (Organic Syntheses, V, 33). The crude hydrocarbon is washed successively with alkali and with water, and is then dried and distilled (yield, 78-87%). cycloHexene reacts with chlorine, bromine, hypochlorous acid, and hydrogen (in presence of Pt or Pd) to give the corresponding chloride, bromide, colorohydrin, and cyclane respectively, and it is oxidised by alkaline permanganate to adipic acid. On long treatment with oxygen it gives a 0.75% yield of the peroxide $C_6H_{10}O_2$, together with a syrup which is probably the dimeric peroxide (Stephens, J. Amer. Chem. Soc. 1928, 50, 568), a much larger yield (20%) of practically pure peroxide is formed when it is irradiated in presence of oxygen (Hock and Schrader, Naturwiss. 1936, 24, 159). This per-Schrader, Naturwiss. 1936, 24, 159). oxide gives, (a) with 2N-sulphuric acid, cyclo-hexanediol, m.p. 104° (50% yield), and cyclopentene aldchyde (10-20% yield), and (b) with concentrated caustic soda, Δ^{1} -cyclohexen-3-ol, which also is produced by reduction of the peroxide with sodium sulphite (Hock and Schrader, *l.c.*). With a 7% or a 20% ethereal solution of peracetic acid cyclohexene yields to the trans glycol, m.p. 103°, by sulphuric acid (Arbusov and Michailov, J. pr. Chem. 1930, [ii], 127, 92); with lead tetracetate it yields 1:2-diacetoxycyclohexane, 3-acetoxy-\Delta-cyclo-bayene as well as 2.2 and 3.6 diacetoxy \Delta 1 hexene, as well as 3:3- and 3:6-diacetoxy-△1cyclohexene (Criegee, Annalen, 1930, 481, 263); with liquid sulphur dioxide in presence of a catalyst (H2O2) it yields a polymeric sulphone (Frederick, Cogan and Marvel (J. Amer. Chem. Soc. 1934, 56, 1815). When its vapour, mixed with carbon dioxide, is passed over finely divided nickel at 180° it undergoes disproportionation to yield a mixture of benzene and cyclobexane:

$$3C_6H_{10} \rightarrow C_6H_6 + 2C_6H_{12}$$

(Böcseken and Sillevis, Proc. K. Akad. Wetensch. Amsterdam, 1913, 16, 499), and when it is treated with aromatic bydrocarbons in presence

cyclene (Bodroux, Compt. rend. 1928, 186, 1005):

$$\begin{split} & \mathsf{C_6H_{10}} + \mathsf{CH_3} \cdot \mathsf{C_6H_5} \to \mathsf{C_6H_{11}} \cdot \mathsf{C_6H_4} \cdot \mathsf{CH_3} \\ & \mathsf{C_6H_{10}} + p. \cdot \mathsf{C_6H_4} (\mathsf{CH_3})_2 \to \mathsf{C_6H_{11}} \cdot \mathsf{C_6H_3} (\mathsf{CH_3})_2 \\ & \mathsf{C_6H_{10}} + 1.3:5 \cdot \mathsf{C_6H_3} (\mathsf{CH_3})_3 \\ & \to \mathsf{C_6H_{11}} \cdot \mathsf{C_6H_2} (\mathsf{CH_3})_3 \end{split}$$

Polymerisation of cyclohexene to the dimeric stage (cyclohexyl-cyclobexene) can be effected with phosphoric oxide (Truffault, Compt. rend. 1935, 200, 406), and to di-, tri-, tetra- and poly- meric stages with boron trifluoride (Hof-

mann, Chem.-Ztg. 1933, 57, 5).

Di- and poly-olefins.—The most important of these are cyclopentadienc, which occurs in the forerun of benzene from coal tar, and the monocyclic terpenes, various of which are eontained in the essential oils of a number of plants. All other members of the group, whether containing five-carbon, six-carbon, or higher rings, are obtained by synthetic means. the exception of cyclopentadiene and the naturally-occurring terpenes (mentbadienes), tbey are almost invariably prepared by the removal of hydrogen halide from dihalogenocyclanes, dihalogeno-cyclenes, or cyclic dibalogeno dienes, either directly by the action of alkalis, e.g. :

cycloHexene dibromide.

$$\begin{array}{c} \text{NaOEt} & \text{H}_2\text{C} & \text{CH} \\ \\ & \text{H}_2\text{C} & \text{CH} \cdot \text{OEt} \\ \\ & \text{CH}_2 & \text{CH} \\ \\ & \text{Heat} & \text{H}_2\text{C} & \text{CH} \\ \\ & \text{H}_2\text{C} & \text{CH} \\ \\ & \text{CH} \\ \\ & \text{CH} \end{array}$$

or more usually by the exhaustive methylation method of Hofmann. According to the latter method the halogeno-compound is treated with an organic base and the resulting amino-compound exhaustively methylated in order to produce a quaternary ammonium salt; this salt, or more usually the corresponding quaternary of anhydrous aluminium chloride, addition of the former occurs at the unsaturated centre of the action of silver oxide or caustic alkali, is distilled, unsaturated hydrocarbon formed, e g ;

It is to be remembered that elimination reactions occurring in a grouping >CH-CHX-CH< may in general take two directions, yielding >C=CH-CH< and >CH-CH=C<, so that mixtures of isomeric dienes and polyenes are frequently obtained by the shove named preparative methods. Such mixtures are usually insepsrable by ordinary fractionation methods, but in these mixtures the conjugated forms appear usually to predominate over and possesses similar additive properties to the unconjugated forms. The physical properties conjugated open-chain dienes. It yields

Δ1 % eyeloHeptadiene

in the forerun has then largely become con- tion of the hydrocarbon, and with quinone

whereon the basic radical is eliminated, and the readily be separated from other materials by fractionation (Perkens and Cruz, J. Amer. Chem. Soc. 1927, 49, 517). The dimeride, containing two isomeric di-cyclopentadienes, is obtained as a low-melting crystalline mass, hp. 56°/13 mm. (Wieland and Bergel, Annalen, 1925, 446, 19; Farmer and Scott, J.C.S. 1929, 177), which progressively reverts on slow refluxing under a short column into monomeric cuclopentadiene The latter again polymerises on atanding and even after a few hours contains a measurable quantity of dimeride, with smaller quantities of higher polymerides. For the structure and stereochemistry of dimeric and trimerie cyclopentadiene, see Ann. Reports, 1932, 29, 113; 1936, 33, 238, cyclo-Pentadiene is especially interesting in that it contains a methylene group, which, owing to its position between doubly-bound carbon atoms is highly reactive. The hydrogen atoms of this methylene group are replaceable by alkali metals, and also react readily with the oxygen of ketones giving rise to coloured hydrocarbon condensation products of the type known as fulzenes, which, like cyclopentadiene, are prons to undergo polymerisation.

cycloPentadiene is a conjugated compound unconjugated forms. The physical properties of the more important cycle dienes, eycle too dibromudes, (i) mp. 45-467, and (ii) and of the more important cycle dienes, and fullers are given in the fall farmer, and fullers are given in the fall farmer and Scott, 1c.), and unites spentances of the fuller form the following the full form the form which has been aged by 460, 93; Alder and Stem, 1cd., 1932, 498; 1917, and the fuller fulle verted into dimeno cyclopentadiene, which can (1 mol. or 2 mols, to 1 mol, of quinone) to give

CONJUGATER COULD, QUENES, AND, POLESNES.

	_	_		_				Rp (°C)	d	π _b
cycloPentadiene								402-416	0 8228°	1-4463182
" Hexadiene .								783-788	0 840420	1.475620
" Heptatriene							ı	116	0 887648 5	1 5213 ¹⁸
" Octatetraene							ļ	30 2-36 4 (14 mm)	0 9254	1 539120
Dimethy Ifulvenc		٠	٠				- }	49-50 (11 mm.)	0 BB1 ₂₀	1 547420
Ethylfulvene .							-	74 (26 mm)	-	-
Methylphenylfulve	ne					-	1	1305 (105 mm)	- 1	-
Diphenylfulvene							. [m p. 82	_	

hy successive elimination of the endomethylene bridges and dehydrogenation into naphthaquinone and anthraquinone respectively.

cycloHexadiene is known in conjugated $(\triangle^{1:3})$ and unconjugated $(\triangle^{1:4})$ forms, which are frequently formed together by dehydrohalogenation of various dibromocyclohexanes or dehydration of dihydroxy-cyclohexanes. Purc \triangle 1:3cyclohexadiene may, however, be obtained by the action of sodium ethoxide on cyclohexene dibromide (Hofmann and Damm, "Mitt. schles. Kohlenforsch. Kaiser-Wilhelm-Ges." 1925, 2, 97). This substance resembles closely the open-chain conjugated diolefins: it differs from cyclopentadiene in that it contains no reactive methylene group and polymerises only on prolonged heating. For the structure of dimerie cyclohexadiene, see Alder and Stein, *l.c.*). $\triangle^{1:3}$ -cycloHexadiene combines taneously with maleic anhydride and with quinone to givo crystalline addition products analogous to those formed from cyclopentadiene. $\triangle^{1:3:6}$ -cycloHeptatriene (Tropilidene)

is a colourless liquid which readily resinifies in air. It has been obtained from cycloheptene dibromide by the Hofmann method (see above), cycloheptadiene heing obtained at an intermediate stage.

tanine (1), the first-formed cyclooctadiene (V) industrial scale an economic proposition.

derivatives (II) and (III) which are convertible | being converted successively into cyclooctatriene and cyclooctatetraene.

cycloOctatetraene is of interest in that despite the fact that its molecule displays unbroken cyclic conjugation, it has no aromatic characteristics; moreover, it resembles in its additive properties and ease of degradation the open-ehain conjugated dienes. Like benzenc, its molecular refraction is normal, exhibiting no exaltation.

CYCLOFORM. Trade name for the isobutyl ester of p-aminobenzoic acid, m.p. 65°. Local anæsthetic.

CYCLONITE (cyclotrimethylenetrinitramine), $(CH_2 \cdot N \cdot NO_2)_3$, also known as "Hexogen" and "T.4," is an explosive which in recent years has been the subject of considerable attention from the military authorities of several countries in the course of the search for more powerful fillings for bombs and other munitions. The compound was first described under the name "Hexogen" in 1899 by G. F. Henning (G.P. 104280), who prepared it by nitrating hexamethylenetetramine, the latter being hexamethylenetctramine, the latter being derived from formaldehyde by condensation with ammonia. The material received little notice, however, until the commercial synthesis cycloOctatetraene has also been prepared of methyl alcohol, leading to cheap supplies of by the Hofmann method from N-methylgrana- formaldehyde, rendered its production on an

500 g. of nature send (sp gr. 1-52) with continuous atirring, the temperature being kept between 20° and 30°C. After the addition of the hexa-methylenetetramine is completed, the mixture is heated slowly to 55°C, at which temperature it is maintained for 5 minutes. The mixture is then cooled to the original temperature and diluted with four times its bulk of water. The precipitated cyclonite is separated, washed first with cold water, then with het dilute soda solution, finally with water again, and then dried. The crude material may be recrystallised from acctone.

This direct nitration of hexamethylene tetmmino has been extensively studied by Ilale (J. Amer. Chem. Soc 1925, 47, 2754), wbo obtained a maximum yield of 75%, using nitric acid of 100% strength and working at a temperature of -20°C.

The industrial production of cyclonite has been developed by the Nobel Dynamito Company of Avighana (Italy) under the name of "T.4"

Details of a process suitable for commercial use are given in B.P. 388615, 1932. According to this method, hexamethylenetetramino (H/M/T) is prepared by bubbling ammonia through a saturated solution of formaldebyde, with suitable cooling. The product is acidified with nitrio acid, thus precipitating the dinitrate of H/M/T, which is filtered off and dried. One part of this salt is then dissolved in 7 parts by weight of very concentrated nitric acid (the actual strength is not stated), the temperature being kept below 15°C for the first stage of nitration. Later, the temperature is stage of miration. Laver, the chaptaness allowed to rise to 30°C, at which it is maintained for about 1 hour. The mixture is then cooled to 0°C, thus precipitating about 60% of the cyclonite, after the separation of which the liquor is distilled at a pressure of a few centimetres of mercury and at a tempera-ture not exceeding 40°C. The nitric acid thus recovered is utilised again in the cycle of operations. The cooling and distillation operaoperations. The cooping and recovering further tions are repeated, thus recovering further amounts of evelonite and nitrie acid. Finally, the liquor is diluted with water, thereby precipitating the remainder of the evelonite dilute acid filtrate is used again for the production of H/M/T dinitrate The filtrate and washings from the latter contain formaldehyde as a secondary reaction product. This is recovered by neutralising the acidity with chalk (or otherwise) and then concentrating and vacuum distilling the liquor. By this procedure the consumption of formaldehydo is reduced to 60% and that of nitric said to 25% of the amounts formerly used, little more than the

theoretical proportions being required.

Physical and Chemical Properties—Crude cyclonite consists of a fine white crystalline powder, which, according to Avogadro (Mém do l'Artillerio Fmnçaise, 1931, 10, 875) melts at 200° to 202°C., with decomposition. The crystals are of the rhembic type, their crystallo graphic constants being described by Terpatra

Manufacture.—According to E. von Hers (Z. Krist, 1926, 64, 169). Large, colourless (U.S.P., 1402993), cyclenite may be prepared on a crystall may be chitained by alsw crystallization from acctone. The melting point of the methylenetetramme in small quantities to precrystallized material is given as 2005 by Ind. Desvergnes (Chim. et 1038). Cyclonite is odourless, tasteless, nonhygroscopie, non-toxic, and unaffected by the setion of light or adverso climatic conditions Being colourless, it has no staming effect on the skin of the workers, nor does it produce any of the usual protant effects common to most other natro-explosives. It is stated that even after breathing appreciable quantities of the material in the form of dust no ill effects were observed. Cyclomite is practically insoluble in cold water (0 007% at 20°C.; 0 15% at 100°C.),

and very sparingly soluble in the common organic solvents, with the exception of acctore, 100 parts of which dissolve 5 parts of cyclonite at 20° and 10 parts at the boiling temperature (57.5°C). The material is also soluble in strong nitrie acid, acetic acid, and hot formic seed From these solutions it is precipitated unchanged on dilution with water. The bulk density of the powder is stated by Avogadro to wary between 08 and 00, the absolute density of the crystals being 1.83, which is higher than that of any other nitro compound (except tetra-nitramline) hitberto used as an explosive. Under high pressure the powder can be compressed to an apparent density of 169 The compound is very stable to heat, no change in weight or appearance being observed after storing for two months at 100°C. or after 48 hours at 120°C. When stabilised by boiling with water and then submitted to the Abel heat test at 80°C., cyclonite produced no coloration on the starch ledido test paper up to 60 hours. On being heated alowly to 202°C, the material melts and begins to decompose, with evolution of gas. If the temperature is raised still further, the decomposition proceeds rapidly to completion. If heated suddenly to 280°C the material explodes violently. When ignited with a flame, bowever, it burns briskly without exploding, the flame having a reddish tinge. Cyclonite is normally regarded as a neutral compound, although according to Henning its solution in acctone has a weakly acid reaction.
The material is unaffected by dilute alkalis,
differing markedly from T.N.T. and similar
intro-explosives in this respect. It is equally resistant to dilute acids and to concentrated hydrochloric acid. It is decomposed, however, by concentrated aulphuric acid with evolution of gas, and is similarly attacked when heated with concentrated alkalis According to Des vergnes it is not possible to estimate cyclonite by the natrometer method owing to the fact that

that the material was reduced quantitatively to C.H.N. by titamum trichloride (Ber. 1921, Explosive Properties.-According to Kast, cyclomic detonates in accordance with the following equation,

54, [H], 318J).

enly about five sixths of the total nitregen is liberated as NO. Rathaburg, however, found

C2H4N4O4-2CO+CO+2H4O+H2+3N5

giving a heat of explosion of 1390 kg.-cal. per kg. | More recent work, however, places this figure at 1,290 kg.-cal., a larger proportion of the carbon remaining as the monoxide than is indicated by the above equation (Nitrocellulose, 1935, 6, 24). Avogadro gives the following constants for cyclonite, the corresponding figures for T.N.T. being shown in brackets by way of comparison: velocity of detonation, 8,380 m. per sec., at density 1.70 (cf. 6,700 m. per sec. for T.N.T. at density 1.58); gas evolved on detonation 908 litres per kg. at N.T.P. (T.N.T. 690 l. per kg.); lead block expansion (Trauzl) test 520 ml. (T.N.T. 310 ml.); compression test (Kast apparatus), 5.2 mm. (T.N.T. 3.6 mm.); figure of insensitivity (Berthe, 2 kg. wt.), 42 cm. (T.N.T. 80, picric acid 60).

Application.—The high velocity of detonation coupled with the large volume of gas liberated and the high density of the original material render cyclonite superior in blasting power and "brisance" to any other explosive hitherto used for bursting charges, not excluding blasting gclatine. In his original patent, Horz suggested that cyclonite might be employed as a bursting charge for projectiles and mines, as a blasting explosive, as a filling for detonating fuze, as an ingredient in percussion caps and detonators, and also as a secondary initiator in fuzes for projectiles. Its use as a propellant is described in G.P. 298539 and G.P. 299028, the following mixture being said to have good ballistic properties: cyclonite 45, guncotton 30, nitrobenzene 10, trinitroanisole 9.4, and tetranitroanisole 5.6%. G.P. 469721, 1927, also described a propellant consisting of cyclonite "phlogmatised" with paraffin or carnauba wax, or with nitroaromatic bodies.

The high melting-point of cyclonite precludes the possibility of pouring it in a molten condition into projectiles or mines. By mixing it, however, with a suitable proportion (from about 30% upwards) of a more easily fusible explosive such as T.N.T., a plastic mixture is obtained which can be poured or screw-filled into the ammunition concerned at a temperature below 100°C. The use of such mixtures also reduces the risk arising from the rather high sensitivity of pure cyclonite without seriously detracting from the superior explosive power of the latter material. In this connection it was observed tbat when a rifle bullet was fired into a bomb filled with pure cyclonite, the latter ignited but did not explode, and the flame soon went out. A stream of bullets from a machine gun, however, caused explosion of the filling (Avogadro, l.c.).

In spito of the difficulties indicated, the future of cyclonite as a major high explosive for military use is regarded as highly promising.

CYCLOPARAFFINS (NAPHTHENES) IN PETROLEUM. Crude petroleum is a mixture of hydrocarbons in which paraffins, cycloparaffins, benzenoid and polynuclear polynuclear hydrocarbons occur. It has been surmised that distillation of petroleum undoubtedly there is a diameter, confusedly aggregated together, or

long series of cycloparaffins, but the separation of naphthenes from paraffins is extremely difficult, even although naphthenes and paraffins can be segregated from other hydrocarbons by the use of specific solvents, as, for example, nitrobenzene and phenols.

In the motor spirit fraction cyclopentane and cyclohexane and their homologues are the main cycloparaffin components. There is evidence that dimethylcyclobutane occurs in Californian petroleum and cycloheptane in Russian oil. The latter hydrocarbon appears to be present in many natural crude oils as does also methylcyclopentane. cycloHexane occurs in Russian gasoline and has been identified in some of the American crudes. Of the other homologues dimethylcyclohexanc, trimethylcyclohexane and dimethylcthylcyclohcxane have obtained from a variety of crudes, e.g. American, Russian, Galician, Borneo Japanese.

In the kcroscne fractions the identification is very much more difficult. Nevertheless, it has been surmised that certain bicyclic hydroaromatic hydrocarbons occur. It is still more difficult to identify hydrocarbons of this type in the higher fractions, e.g. lubricating oils, and, in point of fact, there is very little precise information as to the chemical nature of any of the hydrocarbon components of high boilingpoint.

In the cracking of petroleum cycloparaffins are produced and they conceivably are formed by the cyclisation of preformed elefins. case of cracked Russian kcrosene, methylcyclopentane, cyclohexane, dimethylcyclopentane and methylcyclohexane were identified in the products. So much as 30% of naphthenes have been obtained in an average cracked spirit. Special reference should be made to Carleton Ellis, "The Chemistry of Petroleum Derivatives," Rheinhold & Co.

A. E. D. CYCLORAN E. Potassium oleate+a higher alcohol (b.p. above 160°), used as a wetting-out

CYGNINE, C₁₂H₂₂O₃N₂, is said to have been isolated from *Gastrolobium calycinum* (Leguminosæ of Western Australia) by Mann and Ince (Proc. Roy. Soc. 1907, 79, B, 485). A way of preparing the amorphous alkaloid is described, and also the properties of the free base, the hydrochloride and the chloroaurate, but the report suggests further enquiry.

CYLINDRITE. (Ger. Kylindrit.) A complex sulphide ore of tin, Pb₃FeSn₄Sb₂S₁₄, containing, according to this formula, Sn 25.85%. It forms very peculiar cylindrical rolls of thin folia, the concentric shells parallel to the axis of the cylinder readily separating under pressure. The smooth, bright surfaces of separation no doubt represent the cleavages of cylindrically curved crystals. The colour is blackish leadgrey, with a brilliant metallic lustre. The mineral marks paper, and is difficult to pulverise, cyclopropane and cyclobutane occur in natural in these respects resembling graphite. The ore gas, although the cvidence is not very conclusive. consists of numbers of these rolls, which In the subsequent fractions prepared by the measure up to 3 cm. in length and 1 cm. in

sometimes with a sub parallel grouping, the two name of cymenes. The following is a interspaces being filled with massive material brief account of the more important of these It has been found in some quantity in the Santa hydrocarbons.

Cruz mine, at Poopó, Bohria. Tho above p Methyliso; formula, written in the form

brings out a relation between evandrate and the allied Bolivian minerals, teallite (PhSnS-) and franckette (3PbSnS+Pb FeSb S) Analyses I and II, by G T. Prior (Min. Mag 1904, 14, III, by A. Prenzel (Jahrb Min 1893, 2, 124); IV, E. G. J. Hartley (J.C.S. 1936, 1292),
 V. F. M. Brewer and E. Baker (J.C.S. 1936, 1292).

	1	11	111.	IV	v
Pb.	. 35 24	34 58	35 41	33 97	38 65
Fe	. 281				_
Ag			0 62		
Sň	. 25 65	25.10	26 37	25 06	21 44
Sb	. 1231	1298	8 73	11 92	774
S	. 23 83	23 88	24 50	24 39	
					_
	100 34	99 59	98 63	18 80	91 58
Sp gr.	. 546	5 49	5 42		_

L. J. S. CYMARIN, C₁₀H₁₁O₂, m p 130°, [a]_p +23°, is the simple cymarose glycoside of strephanthidin, i.e. C₂H₁₃O₂ O C₂₃H₂₁O₂ It is present in Canadian hemp and various species of Apocynum. Crystalline Kombé strephanthin is a

mixture of cymana with the glycoside of stro phanthobiose (Jacobs and Hoffmann, J Biol Chem. 1926, 67, 609, 69, 153, 1928, 79, 531) E F. A

CYMAROSE, C,H14O, mp 88°, as the augar obtained from the strophanthidin giveoaide cymarin, present in Canadian hemp as well as in periplocymarin from Periploca graca,

It was shown by Jacobs and Hoffmann (J. Biol. Chem 1926, 67, 603; 69, 153, 1926, 79, 531) to be probably a methyl ether of digi toxose, and ameo it gives riso to hydroxy-methoxyglutaric acid on oxidation with nitric acid (Elderfield, Science, 1935, 81, 440, J Biol Chem. 1935, 111, 527), the position of the methoxy group is located as .

A possible isomeride, sarmentose, has been obtained in crystalline form, m p. 78°, [a], +12°, increasing to +15 8°, from the sarmentory marin

CYMENES (CYMOLS), C10H14 name cymene was first given to the hydrocarbon | Chem. Soc. 1927, 49, 491). Sulphonation gives rolated by Gerhardt and Cahours (Ann Chim the 2 sulphonic acid (Le Flyre, J.CS. 1934, Phys. 1811, [m], 1, 102, 372) from Roman od 1501) of cumin, which is the ethercal oil of Cuminum 2, 5-, 6, and 3 positions, and the higher cyminum. This hydrocarbon was originally chlormation products are accompanied by chlorocalled p-methylpropylbenzene, but Widman toluenes formed by replacement of the isopropyl

p Methylysopropylbenzene (p Cymene),

buted in the essential oils of gymaosperms and Angiosperms, and, more especially, in those of the Umbelisferm and Labiate (Semmler, "Do Aetherischen Oele," iv, 19), Considerable quantities of erude cymene, known as sulphite turpentine, collect on the surface of the liquid in the separator used in recovering sulphur dioxide in the manufacture of wood pulp. Cymeno also occurs in certain petroleums.

Formation and Preparation - Many of the terpenes C10H18 and their derivatives may be readily converted into cymene. Camphor on dehydration yields mixtures of m, and p cymene in proportions varying with the dehydrating agent used, p.Cymene is produced from dipentene when the latter is shaken with concentrated sulphuric acid (Wallach and Brass, Annalen, 1684, 225, 311) and more than 68% of p-oymens is formed when citral is distilled with a httls sodine (Bogert and Fourman, Amer. Perfumer, 1933, 28, 345) The most economical methods of preparing cymene from terpenes are discussed by E Raymond (Bull Soc, chun 1934, [v], 1,

Cymene has been synthesized by many of the classical methods, e.g. from p-brome-cumene (p-bromosspropylbengene), methol include, and sodium in ether (Widman, Ber-ledel, 24, 450), or from isopropyl chlorde, tolueno, and aluminium chlorido (Silver, Bull Soc. chim. 1884, [u], 43, 321). It may be prepared by the action of isopropyl sulphato on magnesium p-tolyl bromido (Bert, Compt. rend 1923, 177, 452; Bull. Soc. chm. 1925, 37, 1252, 1397). It is produced by the alkylation of toluene with propylene or isopropyl alcohol using various catalysts, e g. see J. Amer. Chem See 1935, 57, 893-4, for use of propylene and phosphoric oxide; J. Cen. Chem. Ituss 1935, 5117, for use of isopropyl alcohol and alummeum chloride, Compt. renu. 1935, 200, 466, for use of scopropyl alcohol and sulphunc acid

Propertica -p Cymene is a liquid, b p. 177-177 5° (corr.), d 7° 0 8670 (Perkin, J.C.S. 1896, 69, 1195, 1900, 77, 270), mp. -75°.

Reactions -- For a general study of the chlorinaof Strophanthus sarmentosus (Jacobs and tion, intration and sulphonation of p cyment, Bigelow, J. Biol. Chem. 1932, 28, 355, 449) see T. Kuan [J. Chem. Soc. Japan, 1931, 52, A. 473), p-cymene nitrates successively in the 2-The and 6 positions (Ber. 1887, 21, 2126; J. Amer. Chlorination occurs successively in the (Rich 180), 24, 439, 970, 1623) proved it to group with holorus (Quata all Holorus), Atta be the topropyl compound. Frequently the Acad Alocans, Math. Phys. 1922, 6, No. 14, 30 arous berrene bydroughout of the formula Brome and solo dernatives are known. On C₁₈H₁₈ are grouped together under the collec-

toluic acid, p-a-bydroxyisopropylbenzoic acid, (Jannasch and Fittig, Zeitsch. Chem. 1870, methyl p-tolyl ketone, any of which can be obtained in fair yield under suitable conditions. Catalytic oxidation in the vapour phase over vanadium pentoxide, and in the liquid phase using manganese dioxide have been studied (Ind. Eng. Chem. 1931, 23, 1129; 1932, 24, 1184). In the former case p-toluic acid, and in the latter p-toluic acid and methyl p-tolyl ketone are formed. Cymene is oxidised intestinally to cumic acid. With nitrosyl sulphuric acid in the cold p-cymene undergoes condensation, giving 83% of a dicymene

(Puranen, Ann. Acad. Sci. Fennicæ, 1933, 37A, no. 10).

Tetramethylbenzenes:

telramethylbenzene 1. Consecutive (Prehnitene), C₆H₂(CH₃)₄ (1:2:3:4). When sodium durenesulphonate,

(v. infra), is dissolved in concentrated sulphuric acid and allowed to stand for 12 hours, prehnitenesulphonic acid is formed. The mixture is diluted with water saturated with barium carbonate, the barium salts converted into sodium salts, and these, by treatment with phosphorus pentachloride and ammonia, transformed first into sulphonic chlorides and then into sulphonamides. Prehnitenesulphonamide is separated by means of its sparing solubility in alcobol, and is then hydrolysed by heating with concentrated hydro-chloric acid to 170°, when it yields prehnitene (Jacobsen, Ber. 1886, 19, 1213; see also V. Meyer and Molz, Ber. 1897, 30, 1278). The con-version of durene and isodurenesulphonic acids into prehnitene by the action of strong sulphuric acid is considered in No. VI of a series of papers on the polymethylbonzenes by Smith and Cass (J. Amer. Chem. Soc. 1932, 54, 1614).

Prehnitene may also be obtained by beating

bromopseudocumene,

$$C_6H_2(CH_3)_3Br$$
, $(CH_3,Br,CH_3,CH_3,1:2:3:4)$,

with methyl iodide, benzene and sodium at 150° for 12 hours. It occurs with 1:2:5:6 tetramethylbenzene among the products of dehydrogenation of certain "polyterpinoid" compounds (i.a. see Ruzicka et al., Helv. Chim. Acta, 1934, 17, 426; Naller, J. Amer. Chem. Soc. 1934, 56, 1582

Prehnitene has b.p. 204° and m.p. -4°.

(2) Unsymmetrical tetramethylbenzene (iso-Durene), C₆H₂(CH₃)₄, (1:2:3:5.)—Isodurene may be prepared by the action of dimethyl sulphate on a solution of the Grignard reagent from bromomesitylene (Organic Syntheses, XI, 66). It has b.p. 85-87°/18 mm.

Symmetrical tetramethylbenzene (Durene), C₆H₂(CH₃)₄ (1:2:4:5).—In coal tar (Ber. 1885, 18, 3032). Preparation:—by the action of sodium on a mixture of bromopseudocumene, $C_6H_2(CH_3)_3Br$ (1:2:4:5), methyl iodide

[161]; from toluene and methyl chloride in presence of aluminium chloride (Friedel and Crafts Ann. Chim. Phys. 1884, [vi], 1, 461); from o- or p-xylene, methyl chloride, and aluminium ehloride (Jacobsen, Ber. 1881, 14, 2629). The durenc formed by the two last methods may readily be separated from the accompanying isomeric tetramethylbenzenes on account of its much higher melting-point (Organic Syntheses,

Durene has a campbor-like odour, m.p. 79-80°, b.p. 189-191°. On oxidation with potassium permanganate it is ultimately converted into pyromellitic acid,

 $C_6H_2(CO_2H)_4$ (1:2:4:5).

CYMOLS v. Cymenes.

CYMOPHANE, "oriental cat's eye" v. CHRYSOBERYL.

CYNTHIAXANTHIN. This pigment, which closely resembles zeaxanthin, is associated with an ester of astacene in Cynthia (Halocynthia) papillosa. It separates from alcohol in needles, m.p. 188-189°, and exhibits absorption maxima at 517, 484 and 452 mµ in carbon disulphide (Ledcrer, Compt. rend. Soc. Biol. 1934, 117, 1086).

I. M. H., R. F. P., and F. S. S.

a.CYPERONE.

The sesquiterpene ketone, a-cyperone, $C_{15}H_{22}O$, b.p. 177°/20 mm., d_{25}^{25} 0.9945, $n_{\rm D}^{25}$ 1.5283, $[a]_{5461}$ +138°, $[a]_{5780}$ +118.6°, forms the main constituent of the oil from the tubers of Cyperus rotundus (Hegde and Rao, J.S.C.I. 1935, 54, 387T) and it has been characterised by the preparation of a number of derivatives, the more important being the oxime, m.p. 150.5°, semicarbazone, m.p. 216°, 2:4-dinitrophenylhydrazone, m.p. 208-209°. When digested with alkali or with oxalic acid it yields the storeo-isomerie ketone, β -cyperone, b.p. $175-176^\circ$ / 16 mm., d_{25}^{25} 0.9945, $n_{\rm D}^{25}$ 1.5414, $[\alpha]_{5461}$ +239°, oxime, m.p. 138° , semicarbazone, m.p. 207°, dinitrophenylhydrazone, m.p. 218-219°.

The structure of a cyperone has been

determined by Bradfield, Hegde, Rao, Simonsen and Gillam (J.C.S. 1936, 667) and by Bradfield, Pritchard and Simonsen (ibid. 1937, 760).

The constitutions assigned to a- and β -

cyperones have been confirmed by their syntbcsis (Adamson, McQuillin, Robinson and Simonsen, ibid. 1937, 1576).

L. J. S.

CYPRESS CAMPHOR v. CEDROL. CYPRIAN VITRIOL. Copper sulphate (v. COPPER).

CYRTOLITE, a variety of zircon.

CYSTAZOL

CYSTAZOL, A urinary antiseptic con. J. Biochem. Tolyo, 1928, 8, 441; Toennics and taining hexamethylenetetramine and sodium benzoate. (Allen and Hanburys, London). B.P.C. 1931.

CYSTEINE & CISTINE. CYSTINE. & disulphida-a aminopropionic

HO,C CH(NH,) CH, S S CH, CH(NH,) CO,H,

was first discovered by Wollaston in a urmary calculus (Phil. Trans 1810, 223), Dreched isolated it from the hver of horse (Arch Anat. Physiol 1891, 243) and porpose (Z Biol 1896, 33, 85); it was found by Külz (biol 1890, 27, 415) among the soluble products of an artificial panercatic digestion of fibrm, Morner attinent panetests digested in four assessment aboved it was a product of the hydrolysis of several proteins, being especially abundant in keratins (Z physiol Chem. 1899, 28, 595, 1901, 34, 207) It is now recognised as a constituent of nearly all proteins. Buchtala constituent of nearry au process. Some clibd 1907, 52, 474) obtained the following percentages of cystine from the materials mentioned: human heir, 13-14 5, human nearly, and the constituent of the const 515; horsehair, 798; horse's hoofs, 320; ex hair, 727, hoofs of oxen, 537, pigs' bristles, 7 22, pigs' hoofs, 2-17 | her the systime content of grasses and its relation to wool crop, see Pellard and Chibnall (Biochem J 1934, 28, 326).

For the importance of cystme in metabolism, ace Abderhalden (Z physiol Chem 1915-16, 96, 1-147; Ber 1916, 49, 2449-2473), and Lewis (Physiol. Rev. 1924, 4, 394) In view, however, of the recent discovery of a new aulphur containing amino scid, methionine, much of our knowledge of the physiological rôle of cyatine is under revision (see articles by Leuis and du Vigneaud in Annual Rev Biochem 1933,

1935, end 1936).

When cystine is given with food, it is largely When Cystars is given win 1000, it is a significant ordined to anliheste and excreted as such (Rothers, J. Physiol 1905, 22, 175, du Vigneaud, Craft and Loring, J. Biol Chem. 1931, 104, 81). Baumann and Preusse (Z. physiol. Chem. 1881, 5, 300; Ber. 1885, 18, 258) found that when bromobenzene was administered to onimals it was excreted as bromophenyleysteine,

CO2H CH(NH2) CH2 S C4H4Br,

estheen garethly al gardleters banoquos e or plates, mp. 180-182, and yielding a blue crystallino copper salt, (C,H,BrSNO,) Cu
According to Folin (J. Biol. Chem 1910, 8,

There are various modifications of this method came temperature as the active isomeride, is which differ mainly in the manner of the removal three times as soluble in water, and can be of the free hydrochone acud and the final resolved into the active components by the purification (Henia, J. Biol. Chem. 1011, et al., 2011, 1021, 201

Bennett, J. Biol. Chem. 1936, 112, 39).

The cystine of urmary calculus is identical with that obtained from protein hydrolysis (Rothers, J. Physiol. 1905, 32, 175; Alsberg and Folin, Amer. J. Physiol. 1905, 14, 54; Fischer and Suzuki, Z. physiol Chem. 1905, 45, 405), and was proved to be \$\tilde{\text{P}}\displaysis displays a mino-propionic acid by Predmann (Bettr. Chem. Physiol. Path. 1902, 2, 433) This view of its constitution was confirmed by the synthesis of racemic cystine from ethyl monobenzoylsenie, obtained by reducing ethyl formylhippurate, the product of the condensation of ethyl formate and ethyl hippurate On heating ethyl mono-benzoylserine with phosphorus pentasulphide and hydrolysing the product, cysteine is obtained,

and this is readily converted into cystine by explation with air in an ammoniacal solution;

> H CO,Et+NHBz CH, CO,Et → CHO CH(NHBz) CO,Et →OHCH, CH(NHBz) CO,Et → SH CH, CH(NHB2) CO,Et → SH CH, CH(NH2) CO,H → S CH, CH(NH2) CO,H

S CH, CH(NH,) CO.H

(Erlenmeyer, Ber 1903, 36, 2720) Fischer and Raske (101d. 1908, 41, 893) elso ebtained r cystine from dl a emino \$ chloropropionic acid hydrochloride by heating it with barium hydro-sulphide at 100° in scaled tubes, and oxidising the product by atmospheric oxygen in the presence of ammenia From the natural discrime the cuthers obtained, by a similar process, en optically scive cysine which is identical with the legatine obtained from natural sources. Removal of sulphur by ammoniacal copper solution or by zine gives dl alanine (Mouthner, Z. physiol, Chem, 1912,

Cystine crystallises in thin colourless hexagonal plates, it has no melting point, but decomposes slowly at 258-261° (Neuberg and Majer, Z. physiol Chem. 1905, 44, 472). It is sparingly soluble in water, insoluble in sloohol or acetic acid, soluble in acids or alkalis. The heat of combustion at constant pressure is 4,137 g -cal (Emery and Benedict, Amer, J. Physiol 1911, 28, 301-307). The cystine molecule contains two asymmetric carbon atoms, and laso acid by the addition of solid sodium acetate, a rotation at different p values and found dark heavy precipitate of cystine is obtained, [s]₁₄, -225° at p₀ 01; -325° between p₀ 3 and this is purified by decolorising a solution of 7 and -97° at p₀ 10. (Cystine is converted 3-5%) hydrochlone and with bone black, and to the extent of 50% into resisting by hearing converted to the extent of 50% into resisting by hearing the converted to the cystine is converted to the extent of 50% into resisting by hearing the converted to the cystine is the cystine is converted to the cystine is cystine in the cystine is converted to the cystine is cystine in the cystine in the cystine is cystine in the cystine is cystine in the cystine is cystine in the cystine in the cystine is cystine in the cy precipitating the cystine in the het filtrate by with excess of hydrochloric acid of sp gr. 1-124, hot concentrated sodium acctate solution at 165° for 12-15 hours, this decomposes at the

CYSTINE. 541

isolation of d-cystine (J. Biol. Chem. 1931, 94,

Cystine is readily reduced to cysteine (a-amino-B-thiolpropionic acid), HS.CH, CH(NH2).CO2H (Baumann, Z. physiol. Chem. 1882, 8, 300; Mauthner, Z. Biol. 1901, 42, 176; du Vigncaud, Audrieth, and Loring, J. Amer. Chem. Soc. 1930 52, 4,500; Gebauer-Fülnegg, ibid. 4610). A solution of cystine irradiated with a quartz lamp gives about 5% cysteine (Szendrö, Lampert and Wrede, Z. physiol. Chem. 1933, 222, 16). Cysteine readily yields eystine on oxidation, the change occurring spontaneously in alkaline solution, although the rate is increased by the addition of a mixture of ferric chloride and potassium eyanide (Mathews and Walker, J. Biol. Chem. 1909, 6, 289).

Cysteine and its derivatives are extremely active in facilitating oxidations and reductions in the tissues and it is probable that cystine derives much of its importance in the animal body from the fact that it is a source of cysteine. The most active derivative of cysteine in this respect is glutathione, γ -glutamyl-cysteinylglycine, which has been synthesised by Harington and Mead (Biochem. J. 1935, 29, 1602). This tripeptide has been extensively investigated by Hopkins and co-workers who have shown its occurrence in most tissues in which it promotes oxidation and reduction processes (Biochem. J. 1921, 15, 286; 1925, 19, 787; J. Biol. Chem. 1922, 54, 527; 1929, 84, 269; Proc. Roy. Soc. 1931, B, 109, 58; Biochem. J. 1932, 26, 785, 817; 1935, 29, 108).

Cysteine and cystine can be oxidised by bromine and iodine to cysteic acid (a-amino-β-

sulphopropionic acid),

HO₃S·CH₃·CH(NH₃)·CO₂H,

and from this, by loss of carbon dioxide, taurine, NH₂·CH₂·CH₂·SO₃H, is obtained (Friedmann, Beitr. Chem. Physiol. Path. 1902, 2, 433). According to von Bergmann (ibid. 1903, 4, 92), the normal taurine in the bile originates from the cystine. The oxidation of cystine and cysteine to cysteic acid has been studied in some detail by Shinohara (J. Biol. Chem. 1932, 96, 285) using iodato and by Preisler and Preisler (J. Physical Chem. 1934, 38, 1099) using thallie sulphate. The oxidation with bromine and iodine has also been applied as a method of quantitative estimation (v. infra).

Cystine does not react quantitatively with nitrous acid in van Slyke's method for the estimation of amino-nitrogen (Ber. 1910, 43, 3170). This is probably due to the -SH group being converted in part to sulphurie acid (Lough and Lewis, J. Biol. Chem. 1934, 104, 601).

Cystine and tyrosine can be quantitatively separated by means of absolute alcohol saturated with hydrogen chloride (Plimmer, Biochem. J. 1913, 7, 311). The separation of cystine by means of its copper salt is described by Vickery and Leavenworth (J. Biol. Chem. 1929, 83, 523). They have also investigated the precipitation of eystine as silver salt (ibid. 1930, 86, 129). When eystine in dilute sulphuric acid is treated with silver sulphate and the mixture

for the resolution of inactive cystine and the adjusted to pH 6, 70-85% is deposited, but the precipitate when decomposed with hydrochlorie acid yields cystcine. Accompanying this reduction a small part of the cysteine is oxidised to cystcic acid. The silver compound appears to be a mercaptide. A similar behaviour of cystine with mercuric sulphato has been observed by Andrews and Wyman (ibid. 1930, 87, 427)

and Simonsen (ibid. 1931, 94, 323). For the estimation of cystine soveral methods are available. Harris uses direct isolation (Proc. Roy. Soc. 1923, B, 94, 441) and Vickery and White precipitate cystine as cysteine cuprous mercaptide and estimate the S (J. Biol. Chem. 1933, 99, 701). Rossouw and Wilken-Jorden (Biochem. J. 1935, 29, 219) and Medes and Padis (*ibid.* 1936, 30, 942, 1293) also precipitate as cuprous mercaptide but recover the cysteine and estimate by Folin's method (see below). Volumetric methods for the estimation of eysteine and eystine are based on their oxidation to cysteic acid by bromine and iodine. The reaction is markedly affected by temperature and hydrogen ion concentration (Okuda, A. 1926, 190; 1927, 996; 1929, 730; Lucas and King, Biochem. J. 1932, 26, 2076; Baernstein, J. Biol. Chem. 1930, 89, 124; Virtue and Lewis, ibid. 1934, 104, 415; Lavine, ibid. 1935, 109, The colorimetric methods are all essentially based on colour reactions of cysteine (formed in the case of cystine by reduction with a suitable reagent) and can be adapted for the estimation of cystine, cysteine, or both. That most frequently employed is Folin's and is based upon the blue colour formed by cysteine with phosphotungstio acid. Cystine is reduced by sodium sulphite (Folin and Looney, *ibid*. 1922, 51, 421; 54, 171). The method has been modified in several details (Hunter and Eagles, ibid. 1927, 72, 177; Folin and Marenzi, ibid. 1929, 83, 89; Rimington, Biochem. J. 1930, 24, 1114; Tompsett, ibid. 1931, 25, 2014; Shinohara, J. Biol. Chem. 1935, 109, 665; 110, 263). Another method much used recently is Sullivan's (A. 1926, 1266; 1930, 199, 1604), which depends upon the formation of a red colour when cysteine reacts with 1:2-naphtha-quinone. In the presence of sodium hypo-sulphite the reaction is not given by ergo-thionine or methionine and only by cystine in the presence of a suitable reducing agent. The method has been modified by Lugg (Biochem. J. 1933, 27, 668, 1022) and Bushill, Lampitt, and Baker (ibid. 1934, 28, 1293). The methods of Folin-Looney, Folin,-Marenzi, Sullivan, and Okuda are compared by Sullivan and Hess (A. 1930, 1604).

On heating a solution of eysteine with dimethyl-p-phenylenediamine and ferric chloride . a blue colour is obtained which is claimed to be specific for cysteine and distinct from the methylene blue reaction for sulphides (Fleming, Biochem. J. 1930, 24, 965).

DERIVATIVES.—Cystine forms sparingly soluble salts with copper, C₆H₁₀O₄N₂S₂Cu; or mercury

> C₆H₈O₄N₂S₂Hg₂,HgCl₂; 2C₆H₁₀O₄N₂S₂Hg,HgCl₂,7H₂O; $2C_6H_8O_4N_2S_2Hg_2$, $Hg(NO_3)_2$.

With 30% nitrie or hydrochlorie seid (at room | and by an enzyme in malt extract yielding temperature) it vields compounds

C₄H₁₃O₄N₃S₃·2HNO₄ and C₄H₁₃O₄N₃S₃·2HCl

(Morner, Z. physiol. Chem 1014, 93, 203). Cystine ethyl ester hydrochloride,

S.(CH, CH(NH.) CO, EI]. 3HCL

forms colourless needles, mp. 185° with decomposition (Friedmann, Beite Chem. Physiol. Path. 1902, 3, 1). Gortner and Hoff mann (J. Biol Chem. 1927, 72, 443) giva 181° for the m p of I dibenzoyl eystine and for 1., 170°. They also give the following derivatives of 1 and i cystine; di β naphihalene sulphonyl 1, 203i-cystine: di-p naprinauene suspinary; .
201°; u. 215°, phenylisocyanate l. 145°,
i. 181°, phenylhydanton b. 122-123°, 1,
166°; diethylester dhydrochlorde l. 177°, 1,
185°. I Cystine forms two sparingly soluble addition products with pieric seid, one with 3 mols and the other with 6 mols cystine per miol of pictic acid (Tocnnies, Z. physiol Chem. 1933, 222, 1). Soveral other derivatives are described by Fruton and Clarko (J. Biol. Chem. 1934, 106, 667) Numerous acyl derivatives of cystme are described by Fischer and Suzuki (Z physiol Chem 1905, 45, 405, Tischer and Gerngross, Ber 1909, 42, 1485, Abderhalden and Wybert, Ber. 1916, 49, 2449) Of these di l'ieucyl i cystine,

8:[CH, CH(CO,H)NH CO CH(NH,)C,H,],

is a granular powder, turns yellow at 200°, and decomposes at a higher temperature, has [a]20 -136 6°, nr, when obtained in a crystalline form, line [a] -141 4°, it gives a reddish-violet colour with enpper sulphate, is precipitated by ammonium sulphate, and may he termed an albumore

Mesocystine.-Loring and du Vigneaud (J. Biol Chem. 1933, 102, 287) describe the isolation of the internally compensated stereoisomer of cystine and give proof of its constitu tion. Mesocyatine decomposes at 200-218° diformul. and phenyluramino- derivatives have m p. 192-193° and 191°-192° respectively Loring, Dorfmann and du Vigneaud find that

mesocystino can replace the natural legatine in the diet, d. Cystine does not promote growth. They suggest that mesocystina is reduced in the body with the liberation of l cysteina (J. Biol. Chem. 1933, 163, 399) The cystine isomerides also show differences in the ease with which they are outlised to sulphate in the animal hody (du Vigneaud

Craft, and Loring, ibid 1934, 104, 81)
M. A. W. and W. V. T. CYSTOPURIN. A compound of sodium

acctate and hexamethy lenetetramme (Genatosan, Loughborough). B.P.C. 1931. CYTASE, as recently defined by Oppen heimer, is the group name for the enzymes araban.

75% of xylose (Luers, Woch. Brau. 1928, 45 831

Mannan, galactan and araban are at present ill defined substances, so that little can be said about their appropriate enzymes. Cytases are unlely distributed, but with the exception of tylinase the statements about them are mdefinite (r. Brewino, Vol. Il, p. 87).

CYTISINE. Cytisine is an alkaloid widely distributed in the Fam, Papilionacem; it has been isolated from different plants under various names (ulexine, baptiloxine, sophorine), Plugge and Pluggo and Rauwerds (Arch. Pharm 1895, 233, 430; 1896, 234, 685) have investigated between fifty and a hundred Papilionacem (Cytisus, Ulex, Genista, Sophora, Baptisia, Euchresta, Anagyris, spp), and give the 3 selds they obtained and a complete survey of the occurrence of cytisine. This list is completed by the work of Rauwerda (Chem. Zenit 1998, I, 200). The preids vary con Zenti 1838, i. 2007. Into siene vary cou acherably Cytisus Laburnum (syn Laburnum rulgare) contains roughly 1%; in Sophora scenadifiora it may be as high as 3 47%. In Anagyrus fortida cytisino occurs together with anagyrune (q t.), and Nimethyl cytisine (lng, J C S 1935, 1053) The latter has also been isolated from Caulophyllum thalictroides (Fem. Berbendscew) by Power and Salway (J C.S. 1913, 103, 191) and from Thermopers lanceolata (syn Sophora lupinoides) by Oreeholf, Norkina and Gurewitsch (Ber 1934, 67, [B], 1394).

Extraction (lng. JCS, 1931, 2195, with references concerning earlier extractions)finely ground laburnum seeds (1 kg) are intimately mixed with slaked hmo (100 g) and water (500 c.c) is added; the mixture is then extracted with CHCl, for 20 hours. The solvent is evaporated in rucuo, the residue stirred with light petroleum and left overnight. Most of the alkaloid separates, the mother liquer being extracted with dilute acid. The crude alkaloid is boiled in dilute HCl with animal charcoal, the filtered solution is made strongly alkaline and extracted with CHCls. The duel solution is evaporated, cytisine crystallising out at once (sield 20 g). The alkaloid is best purified by distillation in vacuo, followed by crystalusation from dry acctone,

Gylstene (I), C11H14ON2, [a]17 -119° 57' (in water, for further details concerning rotations, see Rauwerda, J C S. 1900, 78, I, 607; Arch. Pharm 1900, 238, 477); mp. 154 5-155 5°; bp 218°/2 mm.; erystalhses from acctone in prisms. (1) is easily soluble in H₂O, EtOH, CHCl₂, but sparingly soluble in Et₂O and benzene.

N-methyl.(I), [a]D -234° 10' (in H2O), mp 136°, needles (from ligroin or acetone), is obtained by interaction of (1) and methyl toluenesulphonheimer, is the group name for the enzymes ate (Freund and Gauff, Arch. Pharm. 1918, 258, which hydrolyse mannan, galactan, xylan and 33; see also Ing. 1c. 1931). N. methyl (!) methodide, m p. 276° (from MeOH).

Xylan, which Haworth has shown to contain a chain of 18-20 xylopyranosa units, is hydrolysed Pherm. 1872. 230, 4481; mono N benoyl (1) by the enzyme of Heltz, grung 63% of xylose mp. 116° (Mass., Per. 1008, 44, 1635) (Ehrenstein, Helt. Chim. Acta, 1926, 8, 332) tolueneralphonyl (1), mp. 207-208°, soluble in

Treatment of (I) with H₂O₂ yields hydroxy-(I) ,C₁₁H₁₃ON·NOH, m.p. 223-226° (Freund and Friedmann, Ber. 1901, 34, 605). Tetrahydrodeoxy-(I), C₁₁H₂₀N₂, is formed by electrolytic reduction of (I), strongly alkaline oil, b.p. 270°/ 760 mm. (Freund and Horkheimer, Ber. 1906, 39, 814).

In aqueous solution (I) or its salts gives a red colour with ferric salts, which is discharged by H2O2, a blue colour being eventually developed on heating (van de Moer's reaction, see Spath, Monatsh. 1919, 40, 26).

Salts of (I).—(I) is a strong base and forms Salts of (1).—(1) is a strong base and forms crystalline salts, the common salts are monoacidic, but a dihydrochloride has been described (Partheil, l.c., 1892), B·HCl,H₂O, colourless prisms; B·2HCl,3H₂O, yellow needles; B·H1,H₂O, yellow prisms; B₂·H₂SO₄, hygroscopic needles; B·HAuCl₄, reddish-brown needles, m.p. 220°, sparingly soluble in warm H₂O; B·HNO₃,H₂O, needles or leaflets, [2].—81° 90′ $[a]_D - 81^\circ 29'$.

For dissociation constants of (I) and its salts, v. Kolthoff (Biochem. Z. 1925, 162,

Constitution of Cytisine (early references, cf. Ewins, J.C.S. 1913, 103, 97, and Ing., l.c. 1931).— When (I) is heated with HI and red phosphorus to 230°, ammonia, cytisoline (II) (Ĉ₁₁H₁₁ON, needles from EtOH, m.p. 199°), and β-cytisolidine needles from EtOH, m.p. 199°), and \$\beta\$-cytisolidine (III) (\$C_{11}H_{11}N\$, b.p. 132-133°/14 mm., platini-chloride, m.p. 234°) are obtained. (II) and (III) are reduced by sodium and EtOH to a-cytisolidine (IV), (\$C_{11}H_{15}N\$,, oil, platinichloride, m.p. 216°). Zinc dust distillation of (II) affords (III) (Freund, Ber. 1904, 37, 16; Ewins, l.c., 1913). Ewins (l.c. 1913) proved by synthesis (III) to be 6.8-dimethyl guineline. by synthesis (III) to be 6:8-dimethyl-quinoline and (IV) to be 6:8-dimethyl-1:2:3:4-tetrahydroquinoline; (II) is a hydroxy-6:8-dimethyl-quinoline. Spath (Monatsh. 1919, 40, 93) showed that 2-hydroxy-6:8-dimethylquinoline is identical with (II). No further conclusions concerning the constitution of (I) can be drawn from these experiments. Van de Moer's colour reaction (see above) led Spath to conclude (l.c. 1919, p. 26) that (I) contains an α-pyridone group. Ing (l.c. 1931) first demonstrated that (I) does not contain a quinoline nucleus; from theoretical reasons and from the results of a Hofmann degradation he put forward a new formula for (I) which was but slightly modified by Späth and Galinovsky (Ber. 1932, 65, [B], 1526) and accounts satisfactorily for all the degradation products of this compound.

In the Hofmann degradation only the secondary N-atom is involved, the N-atom of the pyridonesystem does not react. Späth and Galinovsky

dilute acid and warm EtOH, insoluble in cold (l.c. 1932) demonstrated that hydrogenation H₂O, can be recrystallised from hot H₂O (Ing., of the des-bases after each interaction of alkali and quaternary base is essential so as to avoid polymerisation. These authors finally obtained tetrahydro-hemicytisylene, C₁₁H₁₅ON (V), which still contains all C-atoms of (I). By ozonolysis of (V), the adjacent pyridone ring is destroyed and a lactam C₇H₁₃ON (VI), formed which may be hydrolysed and then oxidised to a mixture of a-a'-dimethylglutaric acids (VII):

$$(I) \longrightarrow \begin{array}{c} CH_3 \longrightarrow CH \longrightarrow CO \\ CH_2 & N \longrightarrow CO \\ CH_3 \longrightarrow CH \longrightarrow CO \\ V. \\ \\ CH_3 \longrightarrow CH \longrightarrow CO \\ \\ CH_2 & NH \\ \\ CH_3 \longrightarrow CH \longrightarrow CH_2 \\ VI. \\ \\ CH_3 \longrightarrow CH \longrightarrow COOH \\ \\ \longrightarrow CH_2 \\ \\ CH_3 \longrightarrow CH \longrightarrow COOH \\ \\ \end{array}$$

A further important proof for the correctness of formula (I) is provided by Ing (J.C.S. 1932, 2778) and by Späth and Galinovsky (Ber. 1933, 66, 1338). The latter authors effected a Hofmann degradation on N-acetyl-tetrahydrodeoxy-(I), thus opening the reduced pyridone ring and leaving the other heterocyclic ring unattacked. The product obtained was N-acetyl-3-amyl-5methylpiperidine (VIII), which was dehydrogenated and oxidised to B-methylnicotinic acid (IX).

N-acetyl-tetrahydrodeoxy-(I)

$$\begin{array}{ccc} & \text{CH}_2\text{--}\text{CH}\text{--}\text{C}_5\text{H}_{11} \\ & \downarrow & \downarrow \\ & \text{CH}_3\text{CO}\text{--}\text{N} & \text{CH}_2 \\ & \downarrow & \downarrow \\ & \text{CH}_2\text{--}\text{CH}\text{--}\text{CH}_3 \\ & \text{VIII.} \end{array}$$

$$CH_3$$
 C_5H_{11}
 CH_3
 $COOH$
 CH_3
 $COOH$
 $COOH$

(I) is closely related to the alkaloid anagyrine (hands a, b, and c of cytochrome represent the (q t.).

Physiological Action.—(1) is a powerful poison resembling nicotine in its action (cf. Dale and Laid-

CYTISOLIDINE v. CYTISINE. CYTISOLINE t. CYTISINE.

CYTOCHROME. MacMunn in 1886 (Pbil Trans. 177, 267) described a pigment, under the names mychamatin and histohamatin, which he found in muscles and other tissues of most animal species. He concluded that myo- and bisto bamatin are respiratory pigments different from hamoglobin and its derivatives observations were enticised by Hoppe Seyler and others and although MacMunn made a strong defence, his views were not accepted, and his work was forgotten until 1925, when Keilin confirmed his discovery (Proc. Roy. Soc. B. 1925, 98, 312). This worker found that the pigment was more widely distributed and more important than unticipated by MacMuan.

Keilin termed the pigment cytochrome and showed it to be an intracellular substance common to aerobic bacteria, yeast, higher plants and animals. Spectroscopie examination of cells showed it to exist in reduced and oxidised states. In the reduced state it abows, according to Keilin, a characteristic absorption spectrum composed of four bands, a, b, c and d, occupying practically the same position in most of the organisms atudied. In the muscles of guinea-pigs, for instance, the bands are found at (a) 6045Å, (b) 5662Å, (c) 5500Å, (d) 5205Å, whilst the positions of maximum intensity of the bands in the thoracio muscles of bees are (a) 6046Å, (b) 5665Å, (c) 5502Å, (d) 5210Å

The absorption spectrum markedly alters on oxidation, the four etrong bands disappearing and being roplaced by weak diffuse bands lying about a 5665Å and β 5287Å. The highest concentration of the pigment is found mactively metabolising tissues such as the thoracic wing muscles of flying insects, beart muscles of mammals or hirds, pectoral muscles of flying birds or in haker'a yeast.

There is some parallelism between the con-

a bands of the three corresponding compounds. whilst the hand d represents the fused B hands of at least the two compounds b' and c'.

The characteristic properties of cytochrome components are due to the nitrogenous compounds with which their hæmatins are com-bined. "Cytochrome components differ from their hæmatina just as much as hæmoglobin differs from hæmatin" (Keilin). Certain varia. tions in the structure and properties of cyto-chrome are revealed by the study of micro-organisms (Keilin, Nature, 1934, 133, 290). its mining mercinic measure of (f). Lacardo. For which is a shorption spectrum of cytochromes to toxicological identification, see Fuhner and welly mining to that of bakers yeast (a Mertens (Booken Z. 1921, 145, 829); for mero-chemical identification of (f) in plant material, and c may be replaced by a snaple band by see kinen and Farkass (A. 1931, 23). Schl | 5600A), or band a may be shifted a little either towards the violet or the red. It can be definitely stated that the three components of cytochrome "represent three distinct hemochromogen compounds each having at its base a different Fe-porphyrm nucleus It is possible that there nost Fe-porphyrin nucleis are united to similar if not and identical nitrogen compounds. These comment ponents may also differ in the degree of their lies despersion."

Of the three components of cytochrome two. b' and especially a', are very unatable, they are as fragde as most of the enzymes (Keilin, Ergeb. d. Enzymforschung, 1933, II, 239). Component a' decomposes at 55°C, liberating its hamatin compound. Component c' is beat stable and may be easily extracted from baker's yeast, Keilin has atudied the relationship of cytochrome to respiratory enzymes of the cell (Proc. Roy. Soc. 1929, 104, B. 206). His conclusions aro as follows: (1) Components a' and e' of cytochrome are not autoxidisable, whilst b' and an unbound bematin also found in the cell are autoxidisable. The latter in the reduced state can combine with carbon monoxide. (2) The bæmochromogen precursor of cytochrome is also autoxidisable but cannot combine with carbon monoxide. (3) Constituents of cytochrome can give rise to thermostable peroxidase activity. (4) The indophenol oxidase of cells (which is inhibited by cyanide or carbon monoxide) is responsible for the oxidation of cytochrome, especially of its non autoxidisable components a' and c'. (5) Cytochrome in living cells is reduced by organic compounds, especially anceinse acid, when these are activated by their dehydrogenases. (6) Cytochrome acts as a hydrogen carrier between activated organic compounds and activated oxygen (7) The autoxidisable component b' may act aa a carrier between cimilar activated molecules and free molecular ovygen.

Keilm (Proc. Roy. Soc. 1930, 106, B, 418) centration of cytochrome in cells and their has described a method of extracting cyto centration of cytochroms in the prime is chrome of from bakers yeast. Also work and apparently completely absent from anaerous plasmolying Delft yeast with NaCl and pouring into holding tap water. The muture the property of the prime is cooled with its cooled with its property. not one chemically defined substance but a showed to settle, the supernatant fluid removed, mixture of three barmochromogen-like com and the deposit of yeast filtered on a Buchner pounds, a', b', and c', capable of being exidused funnel and washed. The yeast cakes so formed and reduced independently of each other. The are suspended in a solution of sodium hisniphite, sodium hyposulphite (Na₂S₂O₄) is added, and the mixture is left overnight in a stoppered The clear pink filtrate contains, in solution, cytochrome c', mixed with salts and proteins. Subsequent treatment with CaCl₂ and SO₂ effects a separation of the pigment.

The component c' thus extracted corresponds to the modified myohæmatin of MacMunn and shows two strong bands in the visible spectrum, a at 5495Å and β , which seems to consist of three maxima, the central one of which is approximately at 5200Å. Dixon, Hill and Keilin (Proc. Roy. Soc. 1931, 109, B, 29) find that the absolute absorption spectrum of reduced cytochrome c' in the violet belongs to the type of spectrum given by hæmatin compounds. The spectrum of cytochrome c' in the reduced state is that of a typical hæmochromogen and in the oxidised state that of a parahæmatin compound. Reduced cytochrome c' does not oxidise in neutral solution; at $p_{\rm H}4$ and at $p_{\rm H}13$ it becomes, however, autoxidisable. It reacts with carbon monoxide under alkaline conditions, forming a compound easily dissociated by light—behaving thus like an autoxidisable hæmochromogen. "It differs from all known hem and hemochromogen compounds which on shaking with air rapidly oxidise to the corresponding hæmatin and parahæmatin compounds." The constitution of the hæmatin of cytochrome c' is as yet unknown (Roche and Bénévent, Compt. rend. Soc. Biol. 1936, 123, 18). Reduced cytochrome is not oxidised by iron compounds, but by H₂O₂, ferricyanido, and copper salts. Oxidised cytochrome is reduced by sodium hyposulphite, cysteine, p-phonylenediamine, and catechol. Its reduction in the living cell is accomplished by activated hydrogen donators (metabolites), the reduced form being oxidised through the indophenol oxidase. Hence its oxidation in the cell can be inhibited by substances which poison the indophenol oxidase, e.g. cyanide, or H₂S. Under anaerobic conditions cytochrome is found in the living cell in the reduced form which then shows the typical spectrum. Removal of the anaerobic state under normal conditions (e.g. shaking with air or oxygen) causes oxidation of the reduced form of the pigment and the disappearance of the characteristic spectrum. This phenomenon may be observed in living tissues. Recent work (e.g. by Theorell, Biochem. Z. 1935, 279, 463; 1936, 285, 207) has emphasised the important part played by cytochrome as a carrier in the respiration of aerobic cells.

Further information may be sought in an article by Shibata, which deals particularly with bacterial cytochrome (Ergeb. d. Enzymforschung. 1935, IV, 348) and in the papers of Keilin and his colleagues.

J. H. Q. CYTOSINE, 2-oxy-6-amino-dihydropyrimi-

a product of hydrolysis of the nucleic acid from the thyroid gland and of nucleic acids of other origin. It has been synthesised by the following series of reactions: ψ-ethylthio-carbamide hydriodide and ethyl sodioformylacetate condense to form 2-ethylthiol-6-oxydihydropyrimidine

which is converted by the action of phosphorus pentachloride into 2-ethylthiol-6-chloropyrimidine

this reacts with alcoholic ammonia to form the 6-amino-derivative, which yields on prolonged boiling with hydrobromic acid, 2-oxy-6-aminodihydropyrimidine

identical with natural cytosine (Wheeler and Johnson, Amer. Chem. J. 1903, 29, 492, 505; Hilbert and Johnson, ibid. 1930, 52, 1152).

Cytosine crystallises with one molecule of water in thin colourless pearly plates, becomes anhydrous at 100° and decomposes at 320-325°. It is insoluble in ether, and sparingly soluble in alcohol or water, I part dissolving in 129 parts of water at 25°.

DAHL'S ACIDS, I, 2-naphthylamine-5- seeds of Nigella damascena (Linn.) (0.4-0.5%) sulphonio acid; II, 1-naphthylamine-4:6-di and of N. aristata. sulphonic acid; III, 1-naphthylamine-4:7disulphonic acid.

Vor. III.—35

Its constitution was established by synthesis (Ewins, J.C.S. 1912, 101, 544) and "methyl damascenine" (Keller, Arch. Pharm. 1908, DAMASCENINE, methyl 2-methylamino-damascenine" (Keller, Arch. Pharm. 1908, 3-methoxy benzoate, C₁₀H₁₃O₃N, m.p. 25-26°, b.p. 154°/15 mm., 270°/750 mm., occurs in the damasceninic acid, C₀H₁₁O₃N, stated by Pommerehne (16id. 1900, 238, 531) to be isomeric gives the following figures for six samples of with domascenine, is a product of hydrolysis known origin: of the elkaloid (see also Keller and Schulze, sbid. 1925, 263, 481). An alternotive method of synthesis has been described by Kaufmann and Rothlin (Ber. 1916, 49, 578). A full account of the properties and synthesis of demascenine, which is also described as the odoriferona principle of oil of Nigella used in perfumery, is given by Sornet (Manf. Chem. 1924, 5, 87).

Damascenine is mildly narcotic in action. A. J. E.

DAMBONITOL, mp, 195°, is a dimethyl ether of i-inositol; it is found in Gabon rubber and in the latex of Castillon elastica. E F. A.

DAMMAR RESIN. The word used in the Malay States is damar, indicoting resm used for making torches. The word "dammer" is here restricted to the resin used in spirit varnishes ond nitrocellulose lacquers, etc., which is obtained from various species of the Diplero carpacese and possibly Burseracese. The trees producing these resuns are found on the Maleyan Peninsula and the Pacific islands generally. For a full account of the dammar resus obtained in the Federated Malay States and the Straits Settlements, from which the greater part of the world's supply comes, reference should be made to a report by the Conservator of Forests, F.M.S. and S.S. (see J. Oil Col. Chem. Assoc 1924. 7, 186). The resin is obtained by a tapping process, somewhat similar to that used in the production of ordinary turpostine. One of the highest grades of atammar, a name applied to the opaqua rean, sometimes yielded by Balancearpus Cat's Eye dammar, the finest pieces of which Heimi, gave the following figures yield an almost colourless solution in coal tar hydrocarbons. It is mostly obtained from species of Hopea, principally H micrantha The tree exudes resin, which congeals on the trunk. This particular type of tree will not yield a steady supply by tapping. In the Malay Stotes this is known as Mata Kuching From the commercial point of view dammar known as Penck or damar Penck is more important in regord to quantity, although it is not of as high a quality as Mate Kuching. Penak is the examined by F. H. Geake (Ind. Eng. Chem product of Balanocarpus Heimit.

Three samples of dammar examined at the Imperial Institute were found to have the follow- characters: ing choracters.

	2. Brownish- yellow.	
Moisture, % 06	07	1.1
Extreneous matter, % 02	01	28
Acid velue 351	34 6	38 3
Seponification value 396	38 1	478
Ash, % 001	0 01	161
Softening point °C 71 Temp. of graduel	80	72
fusion °C 83-95	88-98	80-90

this resin is shown by the fact that acid values of combustions giving formula such up to 72 have also been found by the Imperial CasHaO a to a resin ocid can hardly be relied Institute.

Coffignier (Bull. S.c. chim. 1911, [iv], 9, 549) (see Dulk, J. pr. Chem. 1848, [i], 45, 16).

20 3 23 6 19 1 4 5 9 6 1 0 224 455 41 379 Insoluble in ether, ... Insoluble in acctone

147 204 140 164 453 218 Origin -1. Padang. 4. Pontlanae 2. Borneo. 3 Singapoi 5. Sumatra. 6. Batyan.

Barry (" Notural Vernish Resins," p. 106 et seq.) gives enalyses of a number of samples which are of considerable value, as the samples were supplied by the Conservator of Forests, F.M.S. and S.S. The samples included Dammer Mata. Kuching, Penek, "dead" dammar, and some inferior dammers. For Mata Kuching be found acid values 21-5, 23 2, and 23 8, and from 0 05-0 47% insoluble in chloroform. For Penak, ocid voluce 34-37 I, and from 0 2-3 84% insoluble in chloroform. The high figures for insoluble matter were found for samples in powder form and may therefore merely indicate high extraneous matter. A sample of Dammar Temak gave the following results:

> Acid value . Saponification value . 25 6 . 1 06% Loss at 105° Melting-point . 80-90 Iodine value (Wns)

	1.	2	3
Moisture (at 105°), %. Insoluble in CHCl ₃ , %. Ash, %. Acid value	0 67	5 86	25
Insoluble in CHCl., %	1.76	Tracea	0-6
Ash. %	0 04	0 12	0.1
Acid value	30 5	26-7	28 4
Non resmous matter, %	2 4	598	3 3
Non resinous matter, % Acid value of pure resin			
resin	31-3	28 4	290

Two samples of Stamese dammars have been 1927, 19, 826), both of which were probably from species of Shorea. They had the following

		1.	2
Specific gravity . Acid value		1 059	1 055
Acid value		32 9	182
Saponification value		159 2	449
Moisture, %		1-9	0.8
Ash, %		0.14	0 35
Saponification value Moisture, % Ash, % Softening-point Melting-point		145°	75°
Melting-point .		193°	85*

Three samples of Stamese resin exemined by Barry (op cut.) gove acid value 17-4-27, saponification value 26 2-45 5, iodine value (Wijs) 43-5-60, insoluble in chloroform 0 4-2 589

It is not considered wise to include the sug-The difficulty in adopting limit figuree for gested composition of this resin, as the results on until crystalline derivatives ere prepared

A so-called Borneo dammar is obtained from | base Vatica moluccana. Kemner (Farben-Ztg. 1936, 41, 586) states that it has acid value 13.47, saponification value 60.37, and melting-point 200°. It contains a little wax, which causes its solutions to be somewhat cloudy. It is known as Hiroe dammar.

E. J. P.

DANAITE. A cobalti-ferrous variety of mispickel (ironsulpharsenide, FeAsS₂), 4-10% cobalt replacing iron.

DANALITE. A rare sulpho-silicate mineral, with the formula $3R_2SiO_4\cdot RS$, where R represents Fe, Zn, Be, Mn, containing BeO about 14%. The flesh-red or grey cubic crystals are octahedral or tetrahedral in habit; sp.gr. 3.35-3.43; H.= $5\frac{1}{2}-6$. It occurs in granite at Rockport and Gloucester, in Massachusetts, and large crystals have been found in Cornwall. Closely allied is the tetrahedral-cubic mineral helvite, which contains about the same amount of beryllium, but with a preponderance of manganese and no zinc.

L. J. S. DANDELION, Taraxacum officinale Wiggers (Fam. Compositæ). A common weed of Europe and America. The leaves, which have a somewhat bitter taste, are sometimes eaten after boiling or used green as a salad. For this latter purpose they are frequently blanched. Recorded analyses of edible leaves are: water, 85.5%; protein, 2.4-2.8%; fat, 0.7-1.0%; N-free extract, 7.5-10.6%; fibre, 1.5%; ash, 2.0%. According to J. S. Kleiner and H. Tauber (Science, 1935, 82, 552) the ascorbic acid content of leaves averages 0.08-0.10 mg. per g. of fresh material. The "down" of dandelion seeds contains, in addition to cellulose, soluble pentoses, hemicellulose and probably a xylan (Votoček and Zvonček Coll Coll Coll (Votoček and Zvoníček, Coll. Czech. Chem. Comm. 1933, 5, 448). Karrer and Schöpp have isolated a flavin from alcoholic extracts of the flowers (Helv. Chim. Acta, 1934, 17, 771).

DAPHNANDRA ALKALOIDS. bark of the Australian plant D. repandula (Fam. Monimiaceæ) is remarkably rich in total alkaloids (6%). Pyman (J.C.S. 1914, 105, 1679) isolated and characterised three new alkaloids from D. micrantha: daphnandrine (I) (1·5–2%), micranthine (II) (max. 0·5%), and daphnoline (III) (1·5%). Mode of extraction: ground bark percolated with 0.25% aqueous tartaric acid, concentrated, made alkaline with NH₃, and extracted with CHCl₃. Separation of phenolic from non-phenolic bases by shaking CHCl₃ extract with NaOH. CHCl₃ evaporated and residue treated with CH₃·CO₂Et, when crude (I) separated. The residue is a mixture of (I) and (II) and is finally separated by conversion into the oxalates. The phenolic portion (see above) is acidified, made alkaline with Na2CO3 and (III) taken up in CHCl3, from which solution the crude phenolic base soon separates.

contains three OMe and one NMe Hydrochloride, B.2HCI+5H2O, groups.

groups. Hydrochloride, B·2HCl+5H₂O, colourless prisms, m.p. 282° (anhydr.), [a]_D +296-314°; hydrobromide B·2HBr+6H₂O, m.p. 291° (anhydr.), [a]_D +265-280°; acid oxalate, B·1½H₂C₂O₄+5½H₂O, m.p. 225°.

Micranthine (II), C₃₆H₃₂O₆N₂, colourless needles from CHCl₃, m.p. 190-196°. The base contains one OMe and one NMe group. Insoluble in H₂O and Et₂O, sparingly soluble in EtOH or CHCl₃, the only crystalline salt is the sulphate B·H₂SO₄+10H₂O, fine colourless needles (from H₂O) decomposing at 312°.

Danholine (III).

Daphnoline (III),

C34H34O6N2 or C35H36O6N2,

crystallises from CHCl₃ or EtOH in small hexahedra with solvent of crystallisation, m.p. $190-215^{\circ}$, $[a]_D + 459^{\circ}$ (dry base in CHCl₃); (III) is even less soluble in organic solvents than (11). Hydrochloride B·2HCl+3½H₂O, large truncated bi-pyramids, from moist EfOH, m.p. 290° (anhydr.), [a]_D +283°. The base contains two OMe and one NMe groups. Hydrobromide B.2HBr+4H2O, clusters of microscopic needles, readily soluble in hot but sparingly so in cold H₂O, m.p. 286° (anhydr.).

The three alkaloids give characteristic colour reactions with Fröhde's reagent: (I) indigo changing to port-wine red; (II) indigo changing to emerald green; (III) violet changing to portwine red.

There is nothing known about the constitution of the three alkaloids, for further colour reactions

and pharmacological action, v. Pyman (l.c.). Schl.

DAPHNETIN. Daphnetin is present in the form of its glucoside, daphnin, in the bark of the Daphne alpina Linn., D. Mezereum Linn., and D, odora (Thunb.).

The concentrated alcoholic extract is digested with boiling water, filtered and treated with lead acetate solution. The precipitate is removed, basic lead acetate added to the filtrate, the lead compound of the glucoside decomposed with hydrogen sulphide, and the solution evaporated (Zwenger, Annalen, 1860, 115, 8). When hydrolysed with boiling dilute acids, daphnin yields daphnetin and glucose.

Daphnetin, C₉H₆O₄, pale yellow needles, m.p. 256°, is soluble in alkalis with a yellow colour, and gives a yellow precipitate with lead acetate solution. It forms a monoethyl ether, colourless plates, m.p. 155°, and a diethyl ether, needles, m.p. 72°. When the latter is boiled with caustic soda solution, the diethyl ether of daphnetinic acid:

is produced, and this, by ethylation, yields the Daphnandrine (I), $C_{36}H_{38}O_{6}N_{2}$, colourless needles from CHCl₃, with 0.5 mol. solvent of crystallisation, m.p. 280°, $[a]_{D}+474.7^{\circ}$ (anhydr.) practically insoluble in boiling H₂O, and all organic solvents except boiling CHCl₃. The

Darhnetin js accordingly 78 dihydroxy-1 coumarin;

(Stünkel, Ber. 1879, 12, 109; Will and Jung, ibid. 1884, 17, 1081).

It has been annthetically prepared by heating pyrogallol with make acid in the presence of sulphuric acid (von Pechmann, thid, 1881, 17, 933). and by the condensation of pyrogalialdchydo with sodium acctate (Gattermann and Koebner, ibid. 1899, 32, 287),

The following derivatives have been prepared . Dibenzoyldaphnetin, mp 152°, diacetyldaph. netin, needles, mp 129° (von Pechmann, Le), m p. 137° (Asai, Acta Phytochem, 1939, 5, 9)

8. Hydrory-7 methorycoumarin, mp. 145°, 8 hydroxy.7 ethoxycoumarin, mp. methozy-7 ethozycoumarin, mp. 81°; and 7-methozy-8 ethozycoumarin, mp. 85°5° (Wessely and Sturm, Ber. 1929, 62, [11], 115; 1930, 63, (B), 1299).

Ethyl daphnetin 3 carboxylate, m p. 231-232° [diacetyl derivative, m p. 129-130°), and the corresponding acid, m p. 263° [diacetyl derivative, m.p. 213-214°) [Bothm, Arch. Pharm. 1033, 271, 490)

Daphnetin 3-carboxylic acid, mp 228°, 8
hydroxy-7-acetonyloxyconmarin, in p. 132-133° and 8 methory-7-acetonyloxycoumarin, m.p 81-82° (Ray, Silooja, and Vaid, J C S 1935, 813) 4. Methyldaphnetin and its derivatives (Sakai

and Kato, J. Pharm Soc Japan, 1935, 55, 691) Daphnotin is a yellon colouring matter and gives very palo yellow, pale chro yellow, chvo-yellow, and chie black shades on nool mordanted with tin, aluminium, chromium and iron respectively (Perkin and Wilson, J C.S. 1003, 83, 134)

Daphnin, CisH10O4, colourless prisms, m.p. 228-229°, dissolves in alkaline solutions with a Sellow colour. Methylation of daplinm and aubsequent removal of the sugar group (Wessely and Sturm, le) yield 7.hudrary-8 methory-coumarin, m p. 1855°. Daplinin, therefore, contains the glucose residue in position 7, and is not identical with the daplinetin gluco-ide synthesised by Leone (Gazzetta, 1025, 55, 673) from 8 acetobromoglucose and daphnetin.

According to Asai (i.e.), the amount of daphnln present in various parts of the D odora varies with the season, e g. large quantities are present in the leaf buds in the early stages of ilevelopment. Daphnin absorbs short waves, especially between the frequencies 3,200 and 3,000, and in this respect is similar to flavone. Coumarin and hydroxycoumarin glucosides resemble the hydroxyflatones in their light absorption and localisation in plants, and probably serve the same physiological purpose. A. G. P. and E. J. C.

DAPHNIN t. DAPHNITING DAPHNOLINE 1. DAPHYANDRA ALKA-

LOIDS. D'ARCET'S ALLOY e. BISMUTH (Vol. I, vinegar, p. 608).

DARCO r. ADSORPTION.

DARWINITE. A copper arsenide, similar to Domeykite, Cu₂As.
DATE. The fruit of the palm Phanix dactylifera. Dates form the historic food of the Arabe, and in recent times have been cultivated in southern areas of the United States. Varieties imported into Europe are usually soft-fieshed and aticky. "Dry," firmer types are more common around desert habitations. Attwater and Bryant (U.S. Dopt. Agric. Off. Lap. Sta. Bull. 1906, No. 28) give the average percentage analysis of the edible flosh of cured ilates as : water 15 4, protein 2-1, fat 2 8, N-free extract + fibre 78 4. ash 13. The soluble carbohydrates of the date consist largely of sucrese, the Deglet Noor (Sahara) variety containing overptionally large proportions. Fattah and Cruess (Plant Physical 1927. 2. 310) record that Mesopotamian varieties contain larger proportions of total solula and of sugars than the American-cultivated types In many varieties the actual sucrose content is higher in the green than in mature fruit, and 20-25% may be inverted before harvesting. Approximately 3% of the carbo-hydrates are in the form of pentosans (Wittmann, Z. landw. Versuelis. Oesterreich, 1991, 4, 131, and according to Reif (Z. Unters. Lebensm. 1931, 66, 179) a small amount of scriptol is also present. A. R. C. Haas and D. E. Blas (Hilgardia, 1935, 9, No. 6, 295) report a detailed study of the growth and composition of the Deglot Noor variety.

Dried ground date stones have the percentage composition: water 7.96, ash 9 80, protein 5 25, carbohydmtes 65 53, fibre 13 60, oil 6 77. The oil has the following characteristics: d15 # 9 9201, 1.1580, saponification value 2061, iodine value 515, unsaponifiable matter 1 98%, free acrd (as oleio) 0 6%, Reichert-Meissl value 10, Polensko value 30, insoluble fatty acids (mp. 22°) 88 7% (R. G. Harry, Analyst, 1936, 61, 403).

The dried, reasted and ground stones have been utilised as a coffee substitute, analysis of which shows :

American German

N-free H.O Protein Fat, extract, Fibre. Ash 53 66 55 70 5001 Includes 2 2% of augur.

The sap of the date palm collected from meisions made in the bark has a high sugar content, and after clarification and evaporation yielda a crudo brown augar or jaggery.

An Indian boverngo propared from dates is stated by Biswas and Guha (Indian Med. Gaz. 1935, 20, 382) to have ten times the vitamin C content of cows' milk, and a considerable proportlen of vitamin-B.

According to Das and Sarin (Ind., Eng. Chem. 1936, 28, 814) vinegar may be antisfactorily prepared from aqueous extracts of dates by inversion of the sugar with acid, and subsequent fermentation; 100 lb. of fresh fruit are stated to yield 12-15 gals, of standard A. G. Po.

This is a native of southern United States and is also grown in the West Indies. The small, round fruits (1.5 cm.) comprise a firm epicarp

enclosing a mass of pulp and seeds.

Pratt and del Rosario (Philippine J. Sci. 1913, 8, 59) record the percentage analysis of the edible pulp and seeds as: total solids, 24.6, insoluble solids 8.4, protein 2.0, acids, as malie, 0.1, reducing sugar 8.1, sucrose 5.3, ash 0.8%.

A. G. Po. DATISCA CANNABINA Linn. This is a tall, erect herb, resembling hemp, belonging to the Datisceæ and met with in the temperate and sub-tropical western Himalayas from Kashmir to Nepal. The root, known as Alkalbir, has been extensively used in Kashmir and throughout the Himalayas as a yellow dyestuff, chiefly on alum-mordanted silk; but the colouring matter appears to be present in the whole plant as the leaves and twigs can also be

employed for the same purpose.

From the leaves of the Datisca cannabina,
Braconnot (Ann. Chim. 1816, [ii], 3, 277)
isolated a crystalline substance which he termed datiscin. It was subsequently shown by Stenhouse (Annalen, 1856, 98, 166) that this compound is a glycoside which on hydrolysis yields a sugar and a yellow colouring matter, datiscetin,

C₁₅H₁₀O₆. Schunck and Marchlewski (Annalen, 1893, 277, 261) suggested the formula C₁₅H₁₂O₆, and showed that, on fusion with alkali, datiscetin gives salicylic acid. By the action of dilute nitric acid, a nitrosalicylic acid was obtained, whereas with strong nitrio acid, picric acid was formed. Datiscetin appeared to contain methoxyl groups, and was considered

by the authors to be most probably a xanthone

derivative.

On the other hand, Korczynski and Marchlewski (Chem. Zentr. 1906, II, 1265; 1907, II, 700; Biochem. Z. 1907, 3, 295) show that pure datiscetin, $C_{15}H_{10}O_6$, does not contain methoxyl groups. It is soluble in alkaline solutions with a yellow colour, and gives with sulphuric acid a yellow liquid possessing a green fluorescence. When boiled with concentrated solutions of alkalis, phenol and salicylic acid are produced.

By the action of bromine on a solution of datiscetin in boiling acetic acid, Leskiewicz and Marchlewski (Ber. 1914, 47, 1599) obtained tribromophloroglucinol, and as a result suggested that this colouring matter is 5:7:2'-trihydroxyflavonol (I). This conclusion was confirmed by

Kalff and Robinson (J.C.S. 1925, 127, 1968), who synthesised datiscetin by the action of o-methoxybenzoic anhydride and sodium omethoxybenzoate on

DATILE. The fruit of Muntingia Calabura. | The datiscetin 3:2'-dimethyl ether thus obtained yielded, on demethylation, datiscotin (I), pale yellow needles, m.p. 276° (corr.), identical with the natural product. From an examination of the actual specimen of datiscotin isolated by Schunck and Marchlewski (l.c.), Kalff and Robinson conclude that the specimen consists essentially of datiscetin mixed with galangin.

Tetracetyldatiscetin, colourless prisms, m.p. 141° (corr.), and tetrabenzoyldatiscetin, colourless needles, m.p. 191-192° (corr.), are described.

Bargellini and Peratoner (Gazzetta, 1919, 49, 64) prepared datiscetin 5:7:2'-trimethyl ether from 2-hydroxy-4:6:2'-trimethoxyphenyl styryl ketone (II), which, by means of dilute alcoholic

hydrochloric acid, yields the corresponding flavanone. The isonitroso- derivative of the latter, when heated with hydrochloric acid in acetic acid solution, is converted into 5:7:2'trimethoxyflavonol, pale yellow needles, m.p. 158-160°, which gives with sulphuric acid an intensely green fluorescent solution.

Datiscin.—According to Charaux (Compt. rend. 1925, 180, 1419), datiscin, C₂₇H₃₀O₁₅,4H₂O, m.p. 192-193°, [a]_D -48·6° in alcohol, occurs to the extent of 6-10% of the dry weight of the roots and leaves of the Datisca cannabina. It is hydrolysed by enzymes yielding equimolecular amounts of datiscetin and a rhamnoglucose, rutinose (cf. ibid. 1924, 178, 1312; Zemplén and Gerecs, Ber. 1935, 68, [B], 1318). By hydrolysis with acids, datiscin gives 1 mol. of datiscetin, 1 of dextrose, and 1 of rhamnosc.

Dyeing Properties of Datisca cannabina.—The dyeing properties of datisca root on wool are very similar to those of Old Fustic. It is, however, slightly inferior in dyeing power. On cotton it dyes more like quereitron bark, in so far that the olives obtained on iron mordant are bluer in shade, as though some tannin matter were present. It appears to have about the same colouring power as quercitron bark, but gives a brighter yellow with aluminium mordant (Hummel and Perkin, J.S.C.I. 1895, 14, 458).

A. G. P. and E. J. C. DATISCETIN AND DATISCIN v. Datisca Cannabina.

DATOLITE. A native borosilicate of calcium, HCaBSiO₅, containing B₂O₃ 21.8%. Monoclinic crystals, with a profusion of small brilliant faces, many with five-sided outlines, are of common occurrence; these are usually transparent and colourless, or with a faint greenish tinge; sp.gr. 3-0. The mineral most greenish tinge; sp.gr. 3.0. often occurs associated with zeolites in the amygdaloidal cavities of basaltic rocks, e.g. in the neighbourhood of Edinburgh and Glasgow, and at Bergen Hill and West Paterson in New ω-methoxyphloraceto- Jersey. It, however, presents several other phenone at 180°, and hydrolysis of the product. modes of occurrence: viz. in granite at Baveno

550 DATOLITE.

in Predmont; in diorite at Rosskopf in Baden; DEAD OIL. The higher boiling fractions of in gabbro with copper ores at Monte Catmi in shale oil, from which the greater parties of the Tuscany; in "limurite" (an aximite-pyroxenite) solid paraffin has been removed by crystallisation at Dundas in Tasmania; in serpentine in the DECACYCLENE, C36H13. formed when Lizard district in Cornwall; in calcute veins in accommendation is heated alone, with sulphus, or sandstone at Sonthofen in Bavaria; in veins of (for greatest yield, 30-35%) with lead oxide in a silver ore at Andreasberg in the Harz Monntains; in the Lake Superior copper mines (here 962; 1914, 47, 1679; 1918, 51, 457), and as a compact, opaque, white form resembling purified by crystallisation from cumene or nitro. unglazed porcelain), in beds of magnetite in gness at Arendal in Norway.

L J. S. DATURA STRAMDNIUM, a hushy annual plant, growing in the warmer temperate regions. The seeds and leaves contain about 0 i to 0 5% of alkaloid (daturine), consisting of hyoscyamine associated with small quantities of

DAURICINE, C36H44N2D4, isolated from Menispermum dauricum (1 3%) by Kondo and

atropine and hyoacine. DATURINE U. DATURA STRAMONIUM.

Nanta (Amer. Chem, Abstr 1927, 21, 2700) Slightly yellow, amorphous hase, mp 115°, [a]0 -139°, soluble in EtOH, acctone, benzene, Et,O. and Dimethiodide needles. /(anhydr) 204°, [a]2 -110°, D.methyldaurseindsmethiodide prisms, m p 152° (decomp), [a]22 -80-1. For absorption spectrum, see Ochiai (Amer. Chem. Abstr 1929, 23, 5272) and Frauendorfer (Ber. 1930, 63, 809) first (Ber. 1913, 46, 2156). suggested that dauricme belongs to the group of the bis coclaumne alkaloids (2 v.) (see also von Bruchhausen and Geruke, Arch Pharm. 1931, 269, 115), which suggestion has finally been proved by the synthesis of methyl daurienne-a-methins, mp. 125-126 (Kondo, Narita, and Uyeo, Ber. 1935, 68, 519, for complete survey, see also Kondo, Arch. Pharm. 1936, 274, 65).

Schl

DEAD DIPPING. The process of producing a pale yellow dead surface on ornmental bethylme e-e-dicathocylle acid has also been breaswork. He breaswork after the final stame of that and the outlaing sweet natural products, ing, with its adhering black scale from the e-g behenoile acid (f. Maruyams, Proc. Imperatural that the reals is detached, when it is removed synthesis has been effected by J. L. Simonem and washed with water. It is next plunged into jand co-workers, J.C.S. 1928, 2673); isodelic acid atronger nitric acid, and when its surface is (K. H. Bauer and J. Mitsotakis, Chem. Umschau, covered with minute gas bubbles, it is washed 1928, 25, 137); secerucio acid (J. L. Simonsen

scaled tube (Dziewoński et al , Ber. 1903, 38, purified by crystallisation from cumene or nitro-benzene. Forms large, glistening, golden yellow needles, mp. 387°, and gives a picrate, mp. 295-296° (dec)

Dziewoński and collaborators have shown by degradation with chromic acid and perman ganate that decacyclene has the constitution :

It may be reduced by hydrodic acid and red ht may be required by symmotr acts and real phosphorus in a sealed tube (Diprovofish et al., J.C.S. 1924, 126, 1, 1056) or catalytically using nickel at 230° and 200 atmospheres pressure (von Braum, Ber. 1934, 67, [B], 214). Sulphonation yields a trisulphonic acid, the alkali salts of which dye wool various shades of red and violet from an acid bath (Dziewoński and Pochwalski, A, 1926, 279, G.P. 379616, USP. 1496085).

Klarner and Woszczerowicz (Chem, Zentr 1934, I, 1760) describe both absorption and

fluorescence spectra.

DECAMETHYLENEOICARBOXYLIC ACID, CD H [CH2]10 CO H. The ester is prepared by electrolysis of potassium ethil pamelate in alkaline solution (G. Komppa, Ber. 1901, 34, 900 . J. Walker and J. Lumaden, J.C.S. 1901, 79, 1200). It is also obtained by adding hydrohromic acid to undecylinic methyl ester and converting this to the nitrile ester with subsequent bydrolysis (Walker and Lumsden, I.c., H. Noerdlinger, Ber. 1890, 23, 2357. For related methods, see J. von Braun, Ber. 1909, 42, 4550; A. Franke and A. Kroups, Monatsh. 1930, 56, 331). A commercial pro-DAVITAMDN. Trade name for prepara-tion containing vitamin A, Dor both Organon Laboratories, London). B.P.C. 1934. Clemmensen method (G.P. 602837). in a solution of argol and dried in bot sawdust and co-workers, J.C.S. 1927, 371); A hydroxy

Pharm. Soc. Japan, 1929, 49, 155); and by reduction of a decadiene acid contained in sardine oil (Y. Toyama and T. Tsuchiya, Bull.

Chem. Soc. Japan, 1935, 10, 192).

Decamethylene-ak-dicarboxylic acid has m.p. 125° (Maruyama), 126.5° (Franke and Kroupa), 129° (corr.) (D. A. Fairweather, Proc. Roy. Soc. Edin. 1925, 46, 71). For X-ray measurements of the acid and esters, see A. Normand, J. Ross, and E. Henderson, ibid. 1926, 47, 69; J.C.S. 1926, 2632.

The half-esters yield esters of eicosane-1:20dicarboxylic acid on electrolysis (D. A. Fairweather, l.c.; L. Ruzicka and co-workers, Helv. Chim. Acta, 1928, 11, 1174). cyclo-Undecanone and cyclodokosane-1:12-dione are formed on distilling the rare earth salts (e.g. thorium and particularly yttrium) of the acid (B.P. 235540; L. Ruzicka and co-workers, Helv. Chim. Acta, 1926, 9, 249; 1928, 11, 670.

DECCAN GRASS CEREALS, (Vol. II, p. 482c).

DECHENITE. Lead vanadite containing zinc and manganese.

DECHOLIN. Trade name for preparations containing dehydrocholic acid, employed in treating hepatic diseases (Riedel- de Haen, Berlin; Old Strand Chemical and Drug Co., London). B.P.C. 1934.

CARBONS DECOLORISING CHARCOAL, ACTIVATED.

DECYL ALCOHOL (and ACETATE). n-Deoyl alcohol, $C_{10}H_{21}$ ·OH, is used, but in very small amount, in perfumes of the rose, neroli, and orange type. Not more than 1% should be used in rose perfumes. Sp.gr. at 15.5° 0.836, $n_{\rm D}^{20}$ 1.4375, b.p. 232°. Decyl acctate, C₁₀H₂₁·OOC·CH₃, has a sweet, fruity odour, and is used to give a distinctive note to many floral perfumes. B.p. 190°.

E. J. P. DECYL ALDEHYDE, n-C, H19. CHO. This synthetic perfume is so powerful that it is usually sold in a 10% alcohol solution, and must be used in traces only. It has an intense, sweet odour, allied to that of orange, and is used—in traces-in floral compounds of the jasmin, rose, violet and wallflower types. When freshly made it melts at 18-20°; b.p. at 12 mm. 94°, at 760 mm. 210°; sp.gr. at 15·5° 0·828; n_D^{20} 1·4296.

DEGALOL. m-Dihydroxycholic acid for the treatment of cholelithiasis. (Riedel-de Haen; Old Strand Chemical and Drug Co.) B.P.C. 1934.

DEGRAS (MOELLON) or SOD OIL is the waste fat recovered as a hy-product from the chamoising (oil-tanning) process, and is used for the currying (stuffing) of heavy leathers and the fat-liquoring of light leathers; its value for these purposes is such that in some factories skins are chamoised solely for the sake of producing, the modllon. For the production of chamois leather, the flesh-splits of sheepskins are used, which are usually de-limed hy "drenching." The wet skins (partially dried by and leath stocking" with sawdust) are stretched on clippings.

stearic acid (Y. Asahina and H. Takimoto, J. | tables and rubbed with fish oil-in England cod (liver)-oil is chiefly used, hut whale oil and scal oil are also employed on the Continent, and menhaden oil in America—and then pummelled and worked in the "stocks" so as to hecome thoroughly impregnated with the oil. The skins are then hung in the air to cool, and the process of ruhbing in the oil, stamping in the stocks and airing is repeated until enough oil has heen adsorhed. By the exposure to the air, the oil undergoes partial oxidation and hydrolysis, which is accompanied by the evolution of volatile products, among which acrolein is noticeable in the later stages. In the so-called English or German methods of working, the oxidation is promoted by piling the skins together in boxes or in covered heaps (after, or between, the stockings) in order to retain the heat generated hy the oxidation; skins must be turned or cooled occasionally, however, in order to prevent overheating. In the French method the skins are "stocked," aired, and "fermented" under milder conditions than in the English method, so that the oil becomes less thickened, and it is possible to recover a large part of the surplus unadsorbed oil by steeping the leathers in warm water and wringing them, or pressing them in hydraulic presses: the fatty matter so expressed is termed première torse moëllon (moëllon dégras in the United States). 1 A further quantity of grease is recovered from the skins by washing them with hot solutions of soda or potash, as is done in the English or German methods; the inferior dégras obtained in this way is frequently added to the moëllon. In the classical English or German method of working, the oil becomes so thickened as a result of the more intense heating that it cannot be pressed out of the skins, and the whole must be recovered by washing with hot alkaline lyes; the emulsion so obtained is acidified with sulphuric acid, when the fatty matter separates and can be skimmed off. This product constitutes the sod oil of commerce. According to Proctor, however, many English manufacturers have now adopted a modified procedure, akin to the French method, and recover a considerable quantity of moëllon by pressure (Proctor, "Principles of Leather Manufacture," 2nd ed., 1922, p. 459); the same is the case in American practice (cf. J. A. Wilson, "Chemistry of Leather Manufacture," A.C.S. Monograph, 1929, vol. II, p. 719).

The mechanism of the changes that occur during' chamoising is still very imperfectly understood. Evidently considerable oxidation and hydrolysis of the oil takes place, and part of the applied unsaturated oil enters into some form of irreversible combination with the skin substance, since only a limited portion of the oil adsorbed by the leather can be recovered by extraction with solvents: moreover, the skin

It may be noted that in the United States wool grease is commonly known as "dégras" or "sod oll," and must not be confused with the "moëllon dégras" obtained from cod-liver oil by chamolsing. In Europe, also, the term "dégras" is Indefinite, as it also embraces the mixed commercial stuffing greases and leather-greases obtained from tanned skins and climping.

to the condition of pelt, as is the case if it is plicated mixture of oxidised (peroxidised, attempted to "chamoise" the skin with a hydroxylated?) acids. saturated acid, e.g. steams acid. Chambard and Michallet (J. Int. Soc. Leather Trades Chem 1927, 11, 559) regard the chamousing process as comprising two distinct reactions, as. (1) the tanning proper, whereby the moist akin aubstance is converted into the more stable leathersubstance by the action of the unsaturated oil with the participation of oxygen (cf. Lauff-mann, Ledertech. Rundschau, 1926, 18, 191), and (2) a secondary "stuffing" or nourishing action whereby relatively inert and polymerised derivatives of the oil are adsorbed by the leather fibrils. The first reaction may take the course suggested by Fahron (Z. angew. Chem. 1891, 4, 172, 634; 1903, 16, 670; 1909, 22, 2083, cf. Meumer and Seyewetz, Coll. 1909, 203, Chim. et Ind. 1918, (1), 79), rez. (1) a peroxida tion of at least two of the unsaturated linkings of the polyethyleme fatty acid chain, followed by (u) the combination (oxidation) of two protein (amino) groups of the skin-substance with one of the peroxide groups, whilst the hydrogen so liberated is absorbed at the second peroxidised linking to form a labile di hydroxy group; the latter may pass finally into a more stable form, perhaps by lactonisation with the carboxyl group liberated by hydrolyais of the oil. Tautomeris change of the dihydroxy groups into the keta hydraxy form, or the conversion of peroxide groups into monoxide (epoxide) groups and other subsidiary oxidation reactions may account for the formation of the inert "oxidised" acids or their glycerides which are found in the unadsorbed mobilion, and for the appearance of other oxidation products (aldehydes, volatile fatty acids, etc.) during the chamosaing process. Fahron's theory that the intermediate, labile oxidation derivatives (peroxides t) of the oils or fatty acids are the actual tanning agents is supported by the fact that neither recovered degras, nor "blown oils," nor the "oxidiaed aculs" from these, possess appreciable chamoising power. The presence of moisture in the skin is essential in order to bring it to the condition of a permeable, re offing to 10 the concentration as a featurement of the control of the probable however, that it are control of the probable however, that the control of the probable however are control of the control is inferior in auppleness and softness to a good chamois leather prepared by means of cod liver oil; a harsh leather is also obtained when fish oil fatty acids are used in place of the glycerides (cf. Fabrion, G.P. 338476; Lewkowitsch and Wood, B.P. 13126, 1911), and the recovered degree is unsatisfactory.

Moëllon possesses considerable emulsifying powers, and contains usually about 20-25% of water; the most characteristic constituents are the resinous "oxidised fatty acids" obtained upon saponification, and termed digras former degragene, Degrasbildner) by Simand (Gerber, 1890, 18, 243, 254, 266, 279); these are insoluble in light petroleum, but soluble in alcohol and alkaline solutions, from which they are re-

undergoes a true tanning ("oil tannage") precipitated by the addition of mineral acid; ance extraction with solvents does not restore it according to Fahrion, they consist of a com-

The characters of pure " moëllon degras " are illustrated by the figures in Table I (due to J. A. Wilson, op. cit., p. 890; Wilson and Herril, "Analysis of Leather," 1931, p. 383); cf. alto Ruhsam, J.S C.I. 1892, 11, 639; Tortelli, Ann. d Lab. chim. d. Gabelle, 1897, 184; Schmitz-Dumont, Dingl. Polyt. J. 1895, 296, No. 9-11; (these, with other figures, are reproduced by Lewkowitsch in "Oils, Fats, and Waxes," vol. III (6th ed.), 1921, pp. 435-9).

Moëllon is not used in its pure state for currying, but is mixed with fish or blubber oils, tallow, etc, and aometimes wool grease, to form the ordinary commercial dégras; according to trade custom, such products are regarded as genume. (Standarda for commercial moèllon gorune. Internative for commercial moellon and degrae, issued by the German Verhand der Degras u. Lederoelfabrikanten (cf. Gnamm, "Die Fettestoffe in der Lederindustria," Stuttgart, 1926, p. 635) are shown in Table II.) On the other hand, numerous substitutes for degras or artificial degras occur in commerce, which consist of highly adulterated degras, the

TABLE I. ANALYSIS OF MOELLON-DEGRAS (WILSON).

	"Typical sample."	Average figures
Free fatty acids (as olcio acid) Unsaponifiable matter Unoxidised fatty acids	21-4 1 2 49 7 (iod. va)	15 7 1-7 51 9 (10d. va)
Oxidised fatty acids .	79; m p 33°C.) 12 8 (iod. val 88)	80; m p 28°C.) 10 4 (iod. val 83)
Water	23·1 1·0	253 08

200 22	ZZAMO.		
	Moellen	"Moellon- Degras,"	Dégran.
Total fatty matter ¹ . Saponifiable matter ¹ . Unexponifiable matter Oxidated acids Volatile matter (water) . Ash	% 80 70 10 6–8 20 ⇒1	78 63 15 5-7 22 >1	75 55 20 4-6 25

DELPHIN.

E. L.

rosin, blown rosin oils, etc.; according to the German standards (see above) rosin is not tolerated in "moëllon" or "dégras," hut may be accepted in products sold as "leather oils"

provided its presence is declared.

For methods of analysis of dégras, cf. Lewkowitsch, "Oils, Fats, and Waxes," vol. III, Ch. 16 (2); Auerbach, Coll. 1931, 311; (provisionally adopted as official methods hy the Int. Ver. Lederind.-Chem.); Immendörfer, Collegium 1925, 135; American standard methods-J. Amer. Leather Chem. Assoc. 1927, 22, 567.

DEGUELIN v. DERRIS.

DEHYDROGENASE. DEHYDRASE, The reducing enzymes are the most labile biocatalysts known. Their isolation from the tissues by methods analogous to those used for the hydrolysing enzymes is usually impossible; perhaps they are closely hound up with the structural elements of the cell. They are usually studied hy Thunberg's (Skand. Archiv. 1920, 20, 1) methylene hlue technique, which consists essentially in determining the reduction time of the dyestuff in the presence of the cell and its substrate, the experiment heing made in vacuum tubes under anaerobic conditions. The substances which reduce the dyestuff are known as hydrogen donators, the substances which can be reduced in presence of the cell are termed hydrogen acceptors and the catalysts as dehydrogenases.

Whilst many substances are active as donators in presence of the cell, many others, closely related in chemical structure, are inert. It is considered, in explanation of this selectivity of action, that there are relatively few dehydrogenases, but that each deals with a particular type of molecule rather than with a specifio substrate (Quastel, Trans. Faraday Soc. 1930, 26, 853). Quastel has emphasised that the essential happening is a change in the substrate molecule as a whole prior to reaction. This is spoken of as the activation of the molecule (Ergebnisse der Enzymforschung, 1932,

209).

Methods of preparing active dehydrogenases from tissues by crushing them very finely in a steel ball mill are described by Ogston and Green (Biochem. J. 1935, 29, 1983, 2005).

The principal dehydrogeneses distinguished act on succinic, malic, formic, citric and lactic

acids, xanthine and alcohol.

Succinic dehydrogenase converts succinic acid into fumario acid in presence of methylene blue. It is present in bacteria and in muscle tissue (Thunherg, Skand. Arch. Physiol. 1917, 35, 165; Wieland and Frage, Annalen, 1929, 477, 1). It has a high specificity.
The so-called Schardinger enzyme of milk

converts aldehyde to acid, hypoxanthin and xanthin to urio acid (Wieland, Ber. 1914, 47, 2085; Annalen, 1929, 477, 32; 1930, 483, 217).

The dehydrase of the acetic hacteria converts alcohol into aldehyde and this into acid, the oxygen acting as acceptor in presence of the first (Wieland, Ber. 1913, 46, 3327; Wieland whon aqueous lead acetate is added gradually

adulterants including mineral oils, blown oils, and Bertho, Annalen, 1928, 467, 95; 1929, 474, 1; Bertho and Baso, Annalen, 1931, 485, 26; also summary hy Bertho, Ergehnisse der Enzymforschung, 1933, 2, 204). The acceptor used for studying these reactions is quinone.

Citric acid dehydrase is present in the liver and in vegetable material acting on citric acid (Bernheim, Biochem. J. 1928, 22, 1178; Thunberg,

Amer. J. Physiol. 1931, 90, 540).

The dehydrases of sceds have been studied hy Fodor (Biochem. Z. 1930, 225, 409, 417; 1932, 246, 4). The subject is too involved for further treatment here; reference is made to the most recent summary articles by Quastel and hy Bertho in Ergehnisse der Enzymforschung, 1933, 1, 209; 1934, 2, 204.

E. F. A.

DEHYDROTHIO.m.XYLIDINE,

is prepared together with ahout an equal amount of isodehydrothio-m-xylidine when m-xylidine is heated with sulphur. The excess of mxylidine is removed by distillation and the residue treated with hydrochloric acid in which the iso-compound is insoluble. Dehydrothio-m-xylidine has m.p. 107°, b.p. 283°/14 mm., is insoluble in water, readily soluble in hot alcohol (Anschütz and Schultz, Ber. 1889, 22, 582; Paul, Z. angew. Chem. 1896, 9, 679), and may be sulphonated (G.P. 593632).

Both dehydrothio-m-xylidine and the isocompound yield azo-dyes, but only those from the latter are substantive to cotton. i80-Dehydrothio-m-xylidine, m.p. 121°, has been shown to have the structure:

(Anschütz and Schultz, Ber. 1925, 8, [B], 64).

DEKOL. Sulphite cellulose waste liquor used as a wetting-out agent.

Trade name for 4:6-dinitro-o-DEKRYSIL. cresol, used in the treatment of obesity (British Colloids, London). B.P.C. 1934.
DELATYNITE v. Amber or Succinite.

DELCOSINE v. DELPHINIUM SPECIES, ALKALOIDS OF.

DELPHIN is the 3:5-diglucoside of delphinidin (q.v.). Qualitative tests (Robinson and Rohinson, Biochem. J. 1931, 25, 1687) had already indicated the possibly wide occurrence of an anthocyanin of this type when Reynolds, Rohinson and Soott-Moncrieff (J.C.S. 1934, 1235) succeeded in isolating delphin from the bright blue flowers of Salvia patens. The pigment, which is present as a complex anyl derivaenzyme. An alternative theory is that a second tive, is accompanied by a large proportion of molecule of aldehyde acts as acceptor for the anthoxanthins. The latter remain in solution 554 DELPHIN.

to a solution of the mixture in methyl alcoholic and s-butyl alcohol. A saturated solution of the HCl, the anthocyanin being precipitated as a instural anthocyanin in 1% methyl alcoholic lead salt. Hydrolysis of acyl groups occurs HClisticapable of dissolving oven fraces of the when the regenerated anthocyann is kept in synthetic sells, and rice sersa; on the other contact with squeous-alcohole HCl, and hand, the chlorides of malvin, cyanin and meconitate with squeous-alcohole HCl. recrystallisation of the simplified pigment gives evanua dissolve in the mixed natural and delphin chloride. The pure diglucoside separates from 3% HCl in homogeneous, flat, leaf-like crystals having a striking hronze reflex, and when dried in air consist of a tribudrate.

The chloride is very sparingly soluble in cold water, alcohol and dilute acids, but it dissolves readily in hot dilute HCI. The pseudo base is formed when the chloride is heated with water or alcohol. The solution in aqueous aodium carbonate is bright pure hive fading slowly to statter and Mieg (Annalen, 1915, 408, 61) by clear green. Sodium acetate gives a violet-blue, hydrolysia of delphinin (q v.), has since been and ferric chloride added to the aqueous solution develops a hlush-violet coloration In alcoholic solution the iron reaction is violet blue A royal hlue precipitate is formed when lead acctate is added to a colution of the chloride. Boiling 10% HCl hydrolyses the anthocyanus to delphinidin chloride (I mol) and glucose (2 mols) Careful comparison of the colour reactions of delphin with those of synthetic anthocyanins of known constitution enabled Reynolds, Rohinson and Scott Moncrieff (l.c.), to relate delphin to cyanin (qv), that is, to assign to delphin the constitution of a 3.5 diglucoside of delphinidin. Delphin chloride was synthesised by condensing together in ice cold ethyl acetate solution saturated with dry HCI 2-O-tetra-acetyl 8glucosidylphloroglucmaldehyde (I) and ietra acetyl. \$\textit{\textit{g}}\ \text{glucosidoxy} - 3 4.5 - tracetoxy - acetophenone (II). An acetylated anthocyann resulted from which delphin chloride (III) was obtained after hydrolysis with methyl alcoholic harrum hydroxide and treatment with dilute sulphurie and hydrochloric acids.

Both the natural and synthetic anthogyanin

synthetic delphin solutious.

Awobanin, the colouring matter of the flowers of Commelina communis var, hortensis Making is a derivative of delphin. Both the chloride and the picrate crystallise with 5H.O. The results of analysis and hydrolysis indicate that it is an ester derived from delphin (1 mol.) and p-commanc acid (1 mol.) (Chika Kuroda, Bull Chem. Soc. Japan, 1936, 11, 265).

DELPHINIDIN, first obtained by Willrecognised as one of the most widely distributed anthocyanidms An acylated monoglucoside, Gentianin, occurs in the flowers of Gentiana acaulus; an acylated rhamnoglucoside, Violania, is present in the flowers of the violet pansy (Viola tricolor L). Vicin, the colouring matter of the scarlet-red vetch (Vicia L), is a mixture of a monoglucoside and a mono rhamnoside of delphinidin. The egg-plant (Solanum Melongena L., var. esculentum Ness) owes its colour to Nasunin, a 3 bioside of delphinidin A dimethyl ether, Maliidin, is a component of many anthocyanina amongst which are Maliin, Oenin (syn. Ampelopen, Cyclamin), Primulin, and Negrelein. A trimetbyl ether, Hirsutidin, occurs in the diglucoside Hirsutin, and Petunin may be the correeponding diglucoside of a delphinidin monomethyl ether (Bell and Robinson, J.C.S 1934, 1604).

Delphiniden chloride, C15H11O,C1,

may be obtained by hydrolysing any of the related glycosides by short boiling with 20% HCl, or synthetically as described below. Willstätter and Weil (Annalen, 1918, 412, 178) have described four distinct hydrates having 1H.O, 1.5H.O, 2H.O, and 4H.O respectively, whilst Reynolds, Robinson and Scott Mouerreff (J.C.S. 1934, 1239) have encountered a fifth form with 3 5H₂O. The monohydrale separates form with 5 bH₂O. The monohydrale separates from cold 3-5% HCl as thin, sharply-cut, deep-violet rhombic tahlets. The sesquihydrale crystallises from HCl of more than 29% concentration. It is readily obtained by adding concentrated HCl to a solution of delphinidin in water or dilute HCI; an amorphous precapitate is first formed but this soon crystallises The dihydrate is obtained when the solvent is allowed to evaporate slowly from a solution of have identical quantitative colour reactions, delphinidin in alcohol to which 7-20% HCl absorption apectra in 0-1% methyl alcoholic has been added; it forms aggregates of prismatic HCl. and distribution ratios between 0.5% HCl tablets. The tetrahudrate separates from solu-

tion in 5% HCl in the form of fine red-hrown | Rohinson, and Schwarzenbach, J.C.S. 1930, prisms and needles. Of the various forms the sesqui- and tetra-hydrates are most easily prepared. Rohinson's hydrate was obtained by dissolving delphinidin chloride in 0.5% HCI and increasing the acid concentration to 10%. The chloride is easily soluble in methyl or ethyl alcohol and in water to violet solutions; it dissolves with difficulty in dilute sulphuric acid. Sodium carhonate added to a solution of the chloride gives a pure hlue coloration. With ferric chloride the alcoholic solution gives a pure blue, whilst in aqueous solution the colour is violet and less stable. The absorption spectrum consists of one fairly well defined hand in the yellow-green region. The anhydrous chloride does not melt helow 350°. The sulphate separates from hot dilute sulphuric acid in long prisms. The picrate forms fine red-brown needles and prisms difficultly soluble in water. The iodide is of particular interest, since it is easily obtained crystalline even when it is derived from crude specimens of delphinidin. It separates when concentrated HI is added to a solution of the chloride in dilute HCl and crystallises from hot dilute HI in elongated rhombic plates which are frequently arranged in rosettes. The crystals are deep-red by transmitted light and exhibit a marked yellow glitter. An aqueous solution of the chloride soon deposits violet flocks of the colour-base. The pseudo-base, C₁₆H₁₂O₈, is formed when a dilute solution of the chloride is warmed with a trace of sodium dihydrogen phosphate. It has considerable crystallising power; extracted with ether and recrystallised from ether or water it forms colourless prisms which become violet when heated above 120°. It is easily soluble in alcohol, acetone, ethyl acetate or glacial acetic acid, but is less soluble in ether and insoluble in benzene. It dissolves in aqueous sodium carbonate to a yellow solution. Dilute HCI regenerates the chloride.

Delphinidin chloride dyes unmordanted wool violet. On tin-mordanted wool the dyeing is blue with a violet tinge. Tannined cotton is

dyed hlue-violet.

An indication of the constitution of delphinidin was ohtained by Willstätter and Mieg (I.c.), who considered that phloroglucinol, pyrogallol, and a small amount of gallic acid were the products obtained on hydrolysing delphinidin chloride with 75% KOH at 250°. Complete proof of structure was provided by Pratt and Rohinson (J.C.S. 1925, 127, 173), who condensed ω -3:4:5tetramethoxyacetophenone with 2-hydroxy-4:6dimethoxybenzaldehyde in ether saturated with dry hydrogen chloride, and ohtained delphinidin chloride hexamethyl ether. Demethylation with HI (sp.gr. 1.7) and phenol gave delphinidin iodide, from which the chloride, identical in all respects with that obtained from natural sources, was obtained by warming in alcoholic solution with silver chloride. A more convenient method of preparing delphinidin consists in condensing 2-O-benzoylphloroglucinal dehyde with ω -3:4:5tetra-acetoxyacetophenone to henzoyldelphinidin chloride, from which delphinidin is obtained by successive treatments in an inert atmosphere with aqueous KOH and HCI (Bradley, tive, and this, after purification, retransformed

793).

Delphinidin chloride may he distinguished from other anthocyanidins in that it is not extracted from 1% hydrochloric acid solution hy the so-called "delphinidin reagent." This consists of anisole (4 vois.) and ethyl amyl ether (1 vol.) containing 5 g. picric acid in 100 c.c. of the mixed solvents. Aerated dilute acid solutions of hoth delphinidin and petunidin are immediately decolorised as the result of oxidation when they are mixed with excess of 10% aqueous sodium hydroxide (R. Rohinson).

DELPHININ, the pigment of the purple wild delphinium (Delphinium Consolida Linn.), was isolated by Willstatter and Mieg (Annalen, 1915, 408, 61). It is an ester of a delphinidin diglucoside (I mol.) and p-hydroxyhenzoic acid (2 mols.), the acyl residues heing attached to phenolic hydroxyl groups (Kondo, Helv. Chim. Acta, 1928, 11, 919). Delphinin is hest isolated as the *chloride*, C₄₁H₃₉O₂₁Cl, which appears to crystallise with 12H₂O, of which only 10H₂O can he removed by drying in high vacuum at 130°. It forms deep red prismatic tablets or prisms, and dried as above sinters at 150-160°, then melts at 200-203° with decomposition and swelling. Although easily soluble in methyl alcohol, it dissolves with difficulty in ethyl alcohol, in HCl of more than 0.5% concentration and in dilute sulphuric acid. The solution in sodium carhonate is hlue, and ferrio chloride added to an aqueous or alcoholic solution gives a blue colour. The salt is optically active; $[a]_{c=}-1364^{\circ}(\pm 150^{\circ})$; $[a]_{614}=-2273^{\circ}(\pm 150^{\circ})$. Its absorption spectrum consists of one broad band. When the neutral aqueous solution is allowed to stand the colour base (rosettes of violet needles from aqueous alcohol) pseudo-base formation is not is formed; observed even when the chloride is warmed with water. The picrate, ohtained as a flocculent red-brown precipitate, is very difficultly soluble in water.

DELPHININE v. DELPHINIUM SPECIES, ALKALOIDS OF.

DELPHINIUM CONSOLIDA. Delphinium Consolida is a common European plant belonging to the larkspur family; its name refers to its powers, real or imaginary, of healing or consolidating wounds. The hlue flowers were examined hy Perkin and Wilkinson (J.C.S. 1902, 81, 585) to determine if these yield the same colouring matters as those previously isolated from the flowers of the D. zalil (ibid. 1898, 73, 267). The presence of kaempferol only, however, could be detected. For its isolation an aqueous extract of the flowers was digested at the boil with addition of sulphuric acid, and the hrown resinous product which separated on keeping, extracted with alcohol, and the extract concentrated. Addition of ether to this solution caused the precipitation of resinous impurity, and on evaporating the ethereal liquid a semi-crystalline residue of the crude colouring matter was obtained. The product was converted into the acetyl derivainto colouring matter. The yield was approxi- with querectin, both apparently as glucosides.

mately 1%. Kaempjerol, C14H10O2, yellow needles, mp. 276-277°, is soluble in alkaline solutions with a yellow colour, and forms crystalline oxonium salts with mineral acids. When fused with alkali, it yields phloroglucinol and p hydroxybenzoic acid.

To keempferol v. Kostanecki (Ber. 1901, 34, 3723) assigned the formula (I), and its synthesis

was aubsequently effected by v. Kostanecki, Lampe and Tambor (såid, 1904, 37, 2096) from 2 hydroxy-4 6 4'-trimethoxychalkone (II) by the usual method of flavonol synthesis, kaempferol 5,7:4'-trimethyl ether, pale yellow needles, m.p. 151-152°, thus obtained yielded kaempferol (I) on demethylation.

Rohmson and Shinoda (J.C S 1925, 127, 1973) aynthesised kaempferol by heating e-methoxyphloracetophenone (III) with anisis anhydride and addium anisate. The product, on hydrolysis, yielded Laempferol 3 4' dimethyl ether, nale yellow platelets, mp. 234°, from which Laempferol was obtained by demethylation

Tetraacetylkaempferol, when cryatalhsed from methyl alcohol, forms colourless needles, and when heated commences to melt at 116° and becomes completely fluid et 120°. On further heating, however, gradual solidification ensues and the product subsequently melts at 181-182°. This peculiarity of acetylksempferol, which is not apparent when the substance is crystallised from ethyl alcohol, affords a convenient method for its detection.

Kaempferol possesses well-defined dyeing properties, and gives with mords need woollen clotb the following shades which closely resemble

those given by morin:

Kaempferol is also present in the Impatiens Balsamina (Chantili Pass), the Erythrina stricta (vernacular name "Kon kathet") (Perkin and Shulman, Proc. Chem. Soc. 1914, 80, 177), the m.p. 1629-163°, is probably a secondary base. berries of the Rhamnus catharticus, and together does not contain methoxyl groups, yields a

in the flowers of the Prunus spinosa (Perkin and

Phipps, J.C.S. 1904, 85, 56). Kaempferide (kaempferol 4'-methyl other) exists in galanga root (q.v.).

The following glucosides of kaempferol are known: Robinin, which occurs in the flowers of Robinia Pseudo-Acacia (q.v.); kaempfentrin, found in the leaves of the Indigofera arrecta (see INDIGO, NATURAL); kaempferin, which has been isolsted from Cassia angustifolia, Vahl. (Tutin, J.C.S. 1913, 103, 2006); multiflorin, present in Rosa multiflora (Konda et al . J. Pharm. Soc. Japan, 1929, 49, 35, 182); and a kaemplerol thamnoside which exists in the blossoms of Acacia discolor, A. linifolia, A. decurrens (var. mollis), and A. longifolia (Petrie, Biochem. J. 1924, 18, 957).

Willstatter and Mieg (Annales, 1915, 408, 61) isolated from the flowers of D. Consolide the

isolated from the anthocyanin, delphinin (q.e.).

A. G. P. end E. J. O. A. KA. SPECIES, OELPHINIUM ALKA-LOIOS OF, Larkspur, pied d'alouette, dauphin (Fr.), Rittersporn (Ger.). Fam, Ranunculacese. About 12 delphinium species are known to contain alkaloids, but only the constituents of D. Ajacis, D. Consolida, and D. Staphisoona have been studied to some extent. Heyl (JCS, 1903, 84, I. 650) isolated a mixture of bases which he called delphocurarine from the root of D. bicolor (0 27%), D. Meniseri (0 35%), D. Nelsonii (0-72%), and D. scoppulorim var. stached, the of the control of the c (1 3%). The bases were extracted by 80% alcohol containing tartane acid. The white amorphous mixture exerted a strong curare like action (d. Lohmann, Pfluger'a Archiv, 1902, 92, 398). From the mixture, a small fraction of a crystalline base, C₁₃H₃₃O,N (!), was isolated, mp 184-185° (from ether and light petroleum), soluble in EtOH, Et.O, CHOI, benzene, eparingly actuble in light petroleum; it contains 18 03% OMe, but does not give crystalline salts or characteristic colour reactions.

Uncharacterised alkaloids are recorded from D. glaucum, D. Geyeri (Heyl, Hepner and Loy, J. Amer. Chem. Soc. 1913, 35, 880), and D Andersons Gray (Miller, Amer. Chem. Abstr. 1923, 17, 3073). According to Keller (Arch. Pharm. 1925, 283, 275), D. elatum contains an alkaloid, C33H51NO, (0 35%), melting at 218°, the only mother liquor is said to contain two more sikalords, but no further details have been reported.

D. Agacis contains alkaloids of a considerably amsiler molecular weight, from an slooholic extract of the seeds Keller and Völker (Arch. Pharm. 1913, 251, 207) succeeded in isolating two new bases

Ajacine, C13H21NO4+H2O, separates from dilute EtOH, m.p. 142-143°, soluble in McOH, EtOH, CHCl₃, benzene, ligroin, scetone, sparingly soluble in Et₃O and CH₃ CO₂Et, insoluble in H₂O. It is alkaline in reaction, yields readily soluble salts, which crystallise with difficulty (bydrochloride, sulphate). base contains three OMe groups.

amorphous di-benzoate.

D. consolida: - (common larkspur, Feld-rittersporn, Ger.). Keller (Arch. Pharm. 1910, 248, 468) isolated three alkaloids from this plant (crude hydrochlorides 1%), one being crystalline, m.p. 195–197°, soluble in EtOH, CHCl₃, acetone, MeOH, sparingly soluble in Et₂O and CH₃·CO₂Et, insoluble in H₂O, strongly alkaline. Markwood (Amer. Chem. Abstr. 1925, 19, 874) re-investigated D. consolida and isolated two crystalline alkaloids, delsoline. C₂₅H₄₁NO₈, m.p. 207-209°, soluble in EtOH and CHCl₃, and delcosine, C₂₁H₃₃NO₆, m.p. 198-199°, soluble in EtOH and CHCl₃, sparingly soluble in Et2O, the latter being probably identical with Keller's crystalline base (see above).

D. Staphisagria: (stavesacre-seeds, Stephanskraut, Läusekraut, Ger.). kraut, Läusekraut, Ger.). For early literature, cf. Walz (Arch. Pharm. 1922, 260, 9). The most important alkaloid is delphinine, C34H47NOg, m.p. $187.5-188.5^{\circ}$, $[a]_{D}^{20}$ + 18.9° , crystalliscs from Et₂O or acetone, soluble in the usual organic solvents. The acid oxalate is characteristic, it crystallises in needles, m.p. 186° (dried). Delphinine is a tertiary base (methiodide, m.p. 196°) it contains one free hydroxyl, one benzoyl, and four methoxylgroups. Delphinine, like most of the D.-alkaloids, is apparently structurally related to aconitine. Keller and Schmechel (Arch. Pharm. 1925, 263, 276) givo a detailed account of the extraction of staphisagria-seeds (yield 0.16% total alkaloids) from which 0.11% crystallised delphinine, small amounts of delphisine, a third crystalline base and amorphous bases were obtained (delphinoidine and staphisagroine (?)). Keller and Schmechel (Arch. Pharm. 1925, 263, 280) further investigated the constitution of delphinine, but so far no definite results have been published. For details concerning the essential oil of D. Staphisagria, cf. Markwood (Amer. Chem. Abstr. 1928, 22, 1215).

Pharmacological Action of Delphinium Alkaloids.—Delphinine is intensely poisonous, resembling aconitine in its action and affecting especially the respiration and circulation by paralysing the nerves of the respiratory system. Lethal dose for dogs per kg. body-weight: delphinine, 1.5 mg., delphisine 0.7 mg., and delphinoidine 0.5 mg. For further details, cf. Lohmann (l.c.), Kahn (Pflüger's Archiv. 1916, 164, 428), and Fühner (Arch. exp. Path. Pharm.

1911, 66, 187).

Schl.

DELPHINOIDINE DELPHINIUM Species, Alkaloids of.

DELPHISINE v. DELPHINIUM SPECIES, ALKALOIDS OF.

DELSOLINE v. DELPHINIUM SPECIES,

ALKALOIDS OF.

DELTA ACID, α-naphthylamine-4:8-disulphonic acid, is a valuable intermediate for the production of 1:8-dihydroxynaphthalene-4-sul-phonic acid, of 5-sulpho-8-amino-a-naphthol, and of a-naphthol-4:8-disulphonic acid. yields a blue wool dyestuff on fusing with sodium sulphide and hydroxide at 210° (G.P. 283727), but is more extensively used in the production of azo-dyestuffs directly 43125) and of tetrakisazo-dyestuffs (F.P. 556489).

crystalline methiodide, m.p. 121°, and an It has also been employed in the preparation of light-sensitive paper which is coated with delta acid and a diazo-component, the red image being

developed with alkali.

DEÑATURANTS. Various articles primarily used for human consumption, such as tea, tobacco, and spirits, are, on importation or manufacture in this country, subject to heavy Customs and Excise duties. When required for other purposes, for example, as raw materials in various arts or manufactures, they are released without payment of duty after having been "denaturing" denatured. This is accomplished by admixture with other substances which, while not impairing their efficiency for the purpose for which they are required, render them unfit for human consumption or act as 'tell-tales" should they be deposited with the Government and "drawback" of duty claimed.

The principles underlying the choice of suitable

denaturants are mainly as follows:

(á) They should be inexpensive so as not to add materially to the cost of the denatured article,

(b) Their use should cause the minimum amount of inconvenience to the manufacturer both in regard to his processes and to tho necessary official supervision involved.

(c) The denaturant should not be incompatible with the process of manufacture or the sub-

sequent use of the denatured article.

(d) If added for its nauseating effect it should be sufficiently objectionable to deter any wouldbe consumers of the denatured article.

(e) If added as a tell-tale it should be distinguishable easily by the microscope or other

(f) It should have such a relation to the original article as to preclude its removal from the mixture by any process which could profitably be carried out on a commercial scale, and it should be capable of being detected even if present in very small quantity.

The over-ruling condition in the use of denaturants is that they shall be approved by the Commissioners of Customs and Excise before

Spirit .- By reason of the large amount of revenue derived from alcoholic spirit and the great variety and extent of its commercial application this is the most important article subject to British official denaturing regulations. There are now in Great Britain four standard types of denatured alcohol, viz.: Industrial Methylated Spirit.

Industrial Methylated Spirit (pyridinised). Mineralised Methylated Spirit.

Power Methylated Spirit.

The quantity of methylated spirit manufactured in Great Britain and Northern Ireland during the past few years has steadily increased. The following table shows the quantity of spirit (expressed in proof gallons) received methylation:

1931-2			10,497,268
1932-3			11,221,933
1933-4		•	12,961,041
1934–5	•		15,416,568
1935-6			16,303,078
1936-7			20,536,112

Industrial methylated spirit, which represents maphthelene or the lime may be replaced by an above quotation, consists of additional quantity of asafoxtida 55 parts by volume of alcohol and 5 parts of Tobacco is almost entirely imported into the wood anghtha (in all methylating operations)
the Commanders of Customs and Excess now process of manufacture coulding. The
have power to approve the use of synthetic
tion of considerable quantities of weste material have power to approve the use or symmetry and the stalks, midn't and proken reals. On the wood distilla-"wood naphtha" in place of the wood distilla-tion product hitherto used exclusively. Indes-duty which has been paid is returned as "draw tion product hitherto used exclusively. Indes-duty which has been paid is returned as "draw to the manufacturer on condition has the calment of the production of the stalks of the same of the stalks of the same official is settredised to composition as industrial methylated spirit either that the tobacco offal is surrendered to composition as managerial integrated sparse, extended to the Covernment or fit intended for each purpose Mineralised methylated sparse coatains 10 parts, as the manufacture of nicotine, meetingles, and 0 5 of crude pyredine. To every 100 gelloms of this maxime is added § agalton of mareral of the maxime is added § agalton of mareral sparse of the maxime is added § agalton of mareral sparse of the maxime is added § agalton of mareral sparse of the maxime is added § agalton of mareral sparse of the maxime is added § agalton of mareral sparse of the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime is a sparse of the maxime in the maxime is a sparse of the maxime is a sparse of the maxime in the maxime is a sparse of the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime is a sparse of the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime in the maxime is a sparse of the maxime naphthe (petroleum od) and not less than 🞝 oz by weight of powdered methyl violet. Power tobacco manufactured in bond (although no methylated spirit is prepared by mixing 100 duty has been paid nor drawback given) in parts by volume of alcohol, not less than 5 parts order to prevent its fraudulent use as a subof petrol or benzol, 0.5 parts of crude pyridine, atitute for tobacco on which duty has been and 25 parts of wood naphtha. To every 100 gellons of this mixture \$\frac{1}{20}\$ oz. of Spirit Red III tobacco to be used in the menufecture of fumidye must be added. When this preparation is gants for horticultural purposes is: mixed with not less than 25% of petrol, benzol or other approved substance, it may be sold free The Statutory of further revenue restriction. Regulations enable the Commissioners of Cus toma and Excuse to permit variations from this prescribed formula It might be mentioned that mineralised methylated epirit, in quantities not exceeding 4 gallons, and the fully denatured lower methylated spirit can be sold by retail With these exceptions a purchaser must have an Exase permit to cover the receipt of these forms of denatured alcohol.

The quantity of spirit used in the manu facture of power methylated spirit is rapidly increasing, as ahown by the following table:

Year.		Proof gallons
1931-2		32,103
1932-3		139,264
1933-4		604,527
1934-5		1,775,519
1935-6		1,556,627
1936-7		5,127,980

The general principles governing the denaturation of alcohol in the Dominions and foreign countries are the same as in Great Britam. variation being made to suit local requirements (v. ALCOHOL).

Tea.—Before being admitted into the United Kingdom, tea is examined by the Customs authorities under Sect. 30 of the Sale of Food and Drugs Act, 1875, and, if found satisfactory, it is passed on payment of duty; otherwise it is thereby separating higher boiling constituents refused edmission as tee for human consumption It still, however, possesses a certain com- heren und Rektifizieren," Leipzig, 1928). mercial value as a raw product for the extraction of theme or caffeine. If it is to be used for this or any other purpose except as a beverage the Commissioners of Customs and Excise perthe Commissioners of Customs and Excess per-physical from bond without paymen and in combination with quinine as of duty on condition that it is suitably de-natured. This is usually accomplished by the addition of 100 parts of line and one of ass-fortiad to 1,00 parts of let. Under special circumstance the assistance where the commissioners, be replaced by the misson of the Commissioners, be replaced Bissurin, Vol. 1, p. 700.

equally necessary in the case of offal from paid. A standard formula for denaturing

Ground tobacco or offal anuff , 92 5 parts 75 Ground moss litter Bone oil 25 .

For certein purposes anthracene oil, in the proportion of 10 paris, is used as an alternative for the bone oil. In these mixtures the oil acts as a deterrent to the use of the preperation as a amoking substitute for tobacco The moss litter is a "tell-tele" to prevent the mixture, after removel of the oil and admixture with other offal snuff, from being fraudulently presented to the Government for drawback. Under certain circumstances the Commusioners of Customs and Excise are prepared to consider applications for permission to use ingredients

supplementary to those prescribed above.
Were which has become unsound and (owing to the formation of acetic acid), end thus unsuitable for use as a beverage, is released from Revenue custody without payment of duty, with a view to conversion into wine vinegar. The denaturant used in this case is 20% of commercial vinegar or an equivelent quantity of acetic acid.

г. с. н. т.

DENDRITES v. AGATE. DENSIMETER. Apparatus for deter-

mining the ap.gr. of liquids DEOXYCARMINIC ACID v. COCHINEAL DEDXYSANTALIN v. BARWOOD

DEPHLEGMATOR. A device for effecting fractional condensation in distillation practice from mixed vapours (K. Thormann, " Destil-

DERBY RED U. AUSTRIAN CINNABAR. DERICINDEL v. CASTOR OIL

DERMATDL, bismuth aubgaliate, is employed as a drug, as a dusting powder (GP. peroxide.

DERMOL. Bismuth chrysophanate,

$Bi(C_{15}H_9O_4)_3Bi_2O_3$ (?)

a yellow amorphous insoluble powder used in the treatment of skin diseases (v. BISMUTH, COMPOUNDS OF, and CHRYSAROBIN).

DERRIS. Derris root has achieved its great importance in Europe and America as an insecticide and rival of pyrethrum since about 1919, prior to which it was known as the source of a fish poison obtained from many species and used by natives throughout the tropics. Although apparently 40 species are known to contain toxio constituents, Derris elliptica, D. malaccensis, and a so-called Sumatra type are the only ones of economic value. Derris root and its extracts (1) are cheaper and more potent than pyrethrum, and different crops are more constant and lasting in action; (2) they are simple and convenient to apply; (3) unlike lead and arsenic preparations which they replace, they are non-poisonous towards higher animals.

The preparation of active extracts consists in extracting with cold or warm water (10 and 16% extracts respectively), with 10% aqueous sodium bisulphite (such extracts are relatively more potent), or best with organic solvents, many industrial processes being based on such extractions. Thus extracts in alcohol or acetone yield stable colloidal solutions on diluting with water; the aqueous emulsion may be concentrated and either precipitated with milk of lime and the solid filtered and decomposed with tartaric acid, or treated with soda and the solid salted out. Powders so obtained have been applied directly, e.g. to tobacco plants, but are more frequently combined with sulphur, soap etc., in the preparation of spraying liquids and cattle washes (B.P. 214822). Other processes merely filter the extract, preferably after adding alcohol and concentrating (such a product contains 30-32% of rotenone and 34-35% of other toxic products) (B.P. 247140) or coagulate with metal salts before re-emulsifying with soaps (B.P. 226250) or sulphonated oils (B.P. 229773). Methyl alcohol, ether, chloroform, oarbon tetrachloride and petroleum are among other solvents which have been suggested for the initial extraction (B.P. 239483). Still other processes employ a conventional protective colloid (gum arabic) to emulsify in water the toxic principles (H. A. Jones and W. M. Davidson, J. Econ. Entomol. 1931, 244) or their solution in higher alcohols (B.P. 285797).

The extraordinary activity of derris preparations is evidenced by the fact that fish are destroyed within 90-240 minutes in a 0.001% aqueous extract (W. Peyer and H. Hünerbein, Apoth.-Ztg. 1931, 92, 1485). Rotenone, the chief toxic constituent, is equivalent at a dilution of 1:100,000 in water to nicotine at 1:10,000 and to pyrethrin at 1:74,800 (M. Darley, J. Econ. Entomol. 1931, 111).

The predominating insecticidal constituents be consulted.

DERMOGENE. Trade name for an anti- of derris root are rolenone, deguelin, tephrosin, septic dusting powder containing 55% of zinc and toxicarol, of relative toxicities 400:40:10:1 (Davidson ibid. 1930, 868). See Derris Resin.

DERRIS RESIN. Derris "root" is the dried rhizome of Derris elliptica, D. malaccensis, and other species. It is indigenous to Malaya and the East Indies. The resin is extraoted from the roots by various organic solvents. It has come into considerable use as an insecticide, and has recently received a great deal of attention from investigators. Its principal toxic constituent is a crystalline body termed rotenone, first isolated by Nagai (J. Tokyo Chem. Soc. 1902, 23, 740). The uncrystallisable portion of the resin, however, is probably as, or nearly as, toxic as the rotenone. Clark heated the uncrystallisable portion with alkali and isolated bodies to which the names toxicarol, deguelin, and tephrosin were given (J. Amer. Chem. Soc. 1930, 52, 2461; 1931, 53, 313, 759; 1932, 54, 2537). These bodies are probably decomposition products of the uncrystallisable resin. Haller and La Forge (ibid. 1934, 56, 2415) obtained dihydrodeguelin by reduction of a fraction of the resin. Cahn and Boam (J.S.C.I. 1935, 54, 42T) have isolated a new substance from what was probably the extract from the root of Derris malaccensis, var. sarawakensis. This substance is dimorphic and melts at 189°, but when crystallised from acetone, melts at 192-194°. It has the formula $C_{23}H_{22}O_{7}$, and is isomeric with tephrosin and toxicarol. It is phenolic and is toxic, but as it is present in very small quantity its presence has little effect on the toxicity. Cahn and Boam consider that the state of knowledge of the constituents of derris resin may be summarised as follows. Rotenone and the new body melting at 189° occur free in the resin; dl-toxicarol and at any rate part of the dl-deguelin do not occur as such. Not more than a small amount of the tephrosin occurs as such. Deguelin occurs, at least partly, either as the l-form or as its simple precursor. The remainder of the deguelin and tephrosin occur in some sort of combination or as simply related precursors. Nothing is known as to the mode of combination of the toxicarol. Cahn and Boam (J.S.C.I. 1935, 54, 37T) consider that the determination of rotenone in derris roots is best effected in the following manner, but if the rotenone content of the resin is below 17% the results are too low.

The root should be powdered to pass a 50-mesh sieve and dried in vacuo before extraction. root is extracted in a Soxhlet tube for 8 hours with trichlorethylene. The solvent is changed and a second extraction for 4 hours is made. If this extraction shows more than a faint yellow colour, a third extraction is made. The solvent is removed and the resin is dissolved in two parts of warm carbon tetrachloride saturated with rotenone. It is cooled, seeded if necessary, and allowed to stand overnight. The separated crystals are collected in a Gooch crucible, washed with CCI4 saturated with rotenone, and dried below 50° in a current of air. The weight of the crystals × 0.72 gives the weight of the rotenone. For fuller details the two papers by Cahn and Boam (l.c.) should

Worsley (J.S.C.I. 1936, 55, 349T) publishes a j very lengthy series of analyses, and considers The sulphonic acids produced in the refining that the following points constitute improve- of beavy fractions of petroleum have found ments in the method of determining rotenure.

Ethyl acetate forms a very efficient solvent for the extraction of the rotenone. The higher the purity of the "rotenone complex" with carbon tetrachloride, the more accurate is the determination, and Worsley adds a given weight of pure rotenone to the extracted remns so as to bring their rotenons value to over 40%. The purity of the carbon tetrachloride com plex prepared by Worsley's method is claumed to be, on the average, 94%, which is much higher than that prepared by Cahn and Boam. There are too many details in the methods for full reproduction here, so that the original papers must be consulted for these.

Tattersfield and Martin (JSCI. 1937, 56, 77T) have separated an optically active coustituent of derris resin which they consider is an active precursor of toxicarol. It is uncertain whether the compound they obtained is in a high state of purity or not. The melting point

was 99° and the specific rotation in 4% benzene

solution was -69° A useful account of the experimental work on the cultivation of Derris elliptica, D. malaccensis, and D. thyrasflora, in Peradeniya, by Lester-James should be consulted (Bull Imp. Inst. 1935, 2, 199).

E J. P DESALGIN. The proprietary name of a preparation containing 0.66% of chloroform in olive oil and peptone stated to have been employed as an analgesic in cases of gastrie dis order (D. Ettinger, Pharm. Presse, 1929, 34, 120).

DESCLOIZITE. A hasse lead and zine vanadate, (Pb,Zn), (OH)VO, crystallising in the orthorhombic system and isomorphous with olivenite (basic copper arsenate). A small amount of copper is usually present replacing zinc, end when this is present in larger amount the mineral grades into cuprodescloizite, which is identical with motiframite and positiacinste (F. A. Banniater, Min. Mag 1933, 23, 376). The cryetals are deep reddish-brown to black in colour, and the streak is a paler yellowish hrown. It occurs in veins of lead ore in association with vanadmute, etc., and has been mined to a himited extent in New Mexico and Arizona, as a source of vanadium. It also occurs in coneiderable quantities in the Sierra de Cordoba in weight values. Aston, using the mass spectro-Argeotina, and at Broken Hill in Northern graph, obtained the value 1-00778±0 00015 Rhodesis. More recently it has been found in for the mass of the hydrogen atom relative to larga amounts at several places in the Groot. D=16 000. This is in agreement with the value fontein district in South-West Africa.

L. J. S DESICHTDL. Trade name for schtbyol which, it is stated, has been deodorised without loss of therapeutic activity by treatment with ateam (Pharm. Zentr. 1898, 39, 90).

OESDXYCINCHONINE v. CINCHONA

CINCHONA ALEALOIDS (this vol., p. 158c).

DESOXYQUININE CINCHONA of 1H.

ALEALORS (this vol., p. 177d).
DESPYRIN. Trade name for a preparasod G. Leemhuis, Apoth .Ztg. 1913, 28, 650).

DETERGENTS FROM PETROLEUM. considerable use as detergents. The sludges contain free acid, polymers, asphalts and a variety of sulphonstes. On addition of water the free acid can be recovered and reconcentrated whilst the only matter separates. From this the sulphonic seids can be recovered by extraction with a number of solvents. Other similar material can be obtained by the sulphonation of paraffin wax and its oxidation products.

Further balogenated sulphonic acids have been proposed. The sulphonic scids thus ohtamed are mixtures of some complexity. They each contain at least one sulphonic and group, but the actual chemical structure of the hydrocarbon associated with it has not been determined. An averaga formula has been suggested of C₂₆H₂₇SD₃H. The sulphonic seid group is very firmly attached to the hydro-carbon nucleus. A larga number of detergent substances are now available from this source. and in general st is the alkali metal calts that have been proposed for this purpose.

There is a considerable patent hat concorned with potroleum detergente, and reference should be made to the very full article in Carleton Ellis's "Chemistry of Petroleum Derivatives,

DETTOL. Halogen derivatives of xylenol dissolved in a saponified mixture of aromatic oils. It is a non irritant and non-toxio germicide

(Reckitt & Sons, Hull). B.P.C. 1934 DEUTERIÚM. introduction. - Denterium "H or D. A heavier isotope of hydrogen having an atomic weight 2-01363±0 00004 relative to helium 4 00216, and 2 01416 relative to oxygen 16 (Bainbridge, Physical Rev. 1933, 44, 57; Wilson, Proc. Roy. Soc. 1936, A, 154, 560). It is present in ordinary electrolytic hydrogen to the extent of 1 part in 30,000± 20% (Bleakney, Physical Rev. 1932, 41, 32; Bleakney and Gould, sbid. 1933, 44, 265; see also Bradley and Urey, ibid. 1932, 40, 889; Unsold, Naturwisa. 1932, 20, 936; Stern and Volmer, Ann. Physik, 1919, 59, 225; Tate and Smith, Physical Rev. 1933, 43, 672; Hardy, Barker, and Dennison, thid, 1932, 42, 279).

History .- The first indication of the existeuce of heavy hydrogen was given by atomic H-1.00780 obtained by chemical methods by

Scott, Morley, Noyes, and Burt and Edgar. After the discovery of the ¹³D isotope and the abundance ratio ¹³D: ¹⁸D, Aston's value calculated to the chemical etandard is H-1.00758, so that the chemical value for ordinary bydrogen is higher by about 1 part in 5,000 than the true value for hydrogen consisting wholly

Barga and Menzel (Physical Rev. 1931, 37, 1669) suggested that this discrepancy was due tion containing 14% of potassium bitartrate to the presence in ordinary hydrogen of a small and 86% of acetylsalcylic acid (C. Mannich amount of a heavier isotope \$H; this fact has been established by subsequent workers Urey,

Brickwedde and Murphy (Physical Rev. 1932, 1 40. 1) set out to concentrate the heavier isotope by evaporation of liquid hydrogen and obtained a residual liquid the Balmer spectrum of which showed clearly a line in the position for an atom of mass 2 and an atomic number 1. Washburn and Urey (Proc. Nat. Acad. Sci. 1932, 18, 496), noted that there was a greater concentration of ²H in the water from old electrolytic cells, and it is this discovery which has enabled the present extensive study of ²H and its compounds to be made. Summaries of the earlier history of this matter are given in the following articles: Nature, 1933, 132, 536; Rutherford, ibid. 1933, 132, 955; 1934, 133, 481.

Much discussion has taken place with regard to the names to be given to 1H, 2H, and possibly 3H. The discoverers suggested protium, deuterium, and tritium respectively (Urey, Murphy and Brickwedde, J. Chem. Physics, 1933, 1, 512), and these names are generally used. Rutherford, however, suggested diplogen for 2H and diplon for the nucleus and the symbol D for the new isotope. Sidgwick supported this view (Proc. Roy. Soc. 1934, 144A, 5). It is generally agreed that "hydrogen" should still be used for the mixture of all the isotopes and that the symbol for 2H shall be D (see Urey, Brickwedde, and Murphy, Nature, 1934, 133, 173).

Occurrence.- Every source of water so far examined shows the presence of heavy hydrogen, but it is difficult to secure an accurate measurement of the amount present. To reconcile the mass spectrograph atomic weight with the chemical value, hydrogen should contain about 1 part of ²H in 4,500. Spectroscopie estimates are much lower, but were suspect owing to the marked tendency of H to be adsorbed in the discharge tube (Lewis and Spedding, Physical Rev. 1933, 43, 964). Later estimates based on mass spectrographic measurements which are probably more reliable put the normal abundance ratio ¹H: ²H at about 1 in 5,000 in rainwater (Bleakney, Gould, *ibid.* 1933, 44, 265); for the method see Bleakney (*ibid.* 1932, 40, 496; also Hardy, Barker and Dennison, *ibid.* 1932, 42, 279; Tate and Smith, ibid. 1933, 43, 672).

Perhaps the greatest obstacle to obtaining a true value of the abundance ratio is due to the large mass ratio (2:1) of the two hydrogen isotopes which, unlike other isotopes, gives rise to sufficient differences in physical and chemical properties as to facilitate partial separation during any process to which ordinary water may be subjected, e.g. distillation, electrolysis, etc. (eee Washburn and Smith, J. Chem. Physics, 1933, 1, 426; Edwards, Bell and Wolfenden, Nature, 1935, 135, 793; Rakshit, J. Physical Chem. 1935, 39, 303; Dole and Wiener, Science, 1935, 81, 45; Christiansen, Crabtree and Laby, Nature, 1935, 135, 870).

Hydrogen occurs for the most part in combination with oxygen as water, and precise determinations of the density of water from various sources have disclosed interesting variations. Water, prepared by the usual methods for conductivity experiments and in addition purified by established methods for density determinations, varied in density over a range

called a gainma of density, $0.000001d = 1\gamma d$), and so is unsuitable as a standard. London Thames water was found to be constant in density and this value agreed with that of surface waters from places so far apart as South Wales, Sumatra, and the U.S.A. Waters from the Dead Sea (3.0yd) and from a Tibetan lake at 13,500 ft. (1.5yd) are heavy, presumably owing to isotopic concentration by evaporation (Briscoe and others, J.C.S. 1934, 1207; 1937, 1492).

The concentration of heavy water in high altitude ice from Alpine glaciers frequently showed an increase, in some instances of as much as 50 to 100 (Baroni and Fink, Monatsh. 1935, 65, 386; 1936, 67, 193; 1937, 71, 128; see also Eucken and Schäfer, Nachr. Ges. Wiss. Göttingen, Math. phys. K1, III, 1935, 1, 137; Riesenfeld and Chang, Ber. 1936, 69, [B], 1305). Variations in the density of rain and snow water from various districts have been observed by Harada and Titani, Bull. Chem. Soc. Japan, 1935, 10, 206, 263; Brodski, Skarre, Donzova and Slutzkaja, J. Physical Chem. Russ. 1937, 10, The water obtained by the combustion of cellulose and a number of other carbohydrates shows an enhanced concentration of deuterium (Harada and Titani, Bull. Chem. Soc. Japan, 1935, 10, 205, 261, 465; Dole, J. Amer. Chcm. Soc. 1936, 58, 580).

The general conclusion from observations on the enrichment of deuterium oxide in waters obtained from natural sources is that such enrichment is comparatively small and is attributable to the operation of physical processes, e.g. preferential evaporation, freezing or diffusion.

Separation of the Isotopes.—(a) Electrolytic Concentration of Deuterium Oxide.-The production of large amounts of pure deuterium oxide is important from the research worker's point of view, and its possible technical applica-tions have still to be explored. The most suitable process appears to be the electrolytic enrichment of ordinary water, and much work has been done in this direction in a relatively short time. Washburn and Urey (Proc. Nat. Acad. Sci. 1932, 18, 496) showed that when dilute sulphuric acid is electrolysed using platinum electrodes a residual water having an increase in density of 164yd was obtained after 98% of the water was decomposed. Further work was done by Lewis and MacDonald (J. Chem. Phys. 1933, 1, 288, 341, 481); Washburn, Smith and Fransden (Bur. Stand. J. Res. 1933, 2, 453; J. Chem. Phys. 1933, 1, 288); Harkins and Doede (J. Amer. Chem. Soc. 1933, 55, 4330); Harteck (Proc. Physical Soc. 1934, 46, 277; Proc. Roy. Soc. 1934, 144A, 1); Newell and Ficklen (J. Amer. Chem. Soc. 1933, 55, 2167); Anderson, Holford and Bates (J. Chem. Phys. 1934, 2, 342). Taylor, Eyring and Frost (ibid. 1933, 1, 823), obtained 82 c.c. of 99% D₂O from 610 gallons of electrolytic liquor. This method with slight variations is the one generally employed in industry. 0.5 N-sodium hydroxide is electrolysed between nickel or iron electrodes, the electrolyte being kept below 20°C. The process is carried out in about seven stages. In the first stage electrolysis proceeds until the volume is reduced of 5.4 γd (one-millionth of the ordinary unit is to about 1/10, when the electrolyte has become

concentrated in alkali and the process is stopped. About 9/10 of the electrolyte is for the separation of neon isotopes, prepared treated with CO., ilistilled to drypess and the distillate added to the remaining 1/10 of the [Naturwiss, 1933, 21, 884; eee also MacGillavry, electrolyte, thus regaining approximately the Rec, trav. chim. 1937, 56, 330). See DIFFUSION, electrolyte, thus regaining approximately the original alkali conventration. This process of electrolysis and removal of alkalı is repeated until the water has reached the desired degree of heaviness. Taylor, Eyring and Frost's results (l.c.) are given in the following table:

ter obtained from lectrolysis No.	Density die	% H3 droger which is bear
I	0 996	_
II	0 333	0.5
III	1 001	25
IV	1 007	8
V	1 031	30
VI	1 098	93
VII	1 104	99

aulphuric acid and lead electrodes is described by Tronstad and Brun (Z. Elektrochem 1934, 40, Gartner (Helv. Chim Acta, 1934, 17, 970, 556). The efficiency of the concentration is 1008), Jaulmes (Chim. Ind. 1935, 33, 1045); remarkably insensitive to the conditions of Stedman (Canad. J. Res. 1935, 13B, 114); electrolysis, and, although the separation factor [Hall and Jones (J. Amer. Chem. Soc. 1934, 56, has been the subject of extensive work, nothing [749] Chemical methods were tried by Davis aufficiently definits to have real significance has and Johnston (J. Amer Chem. Soc. 1934, 56, so far been forthcoming Under apparently 492), Highes, Ingold and Wilson (J.C.S. 1934. identical conditions different results have been [493] obtained (cf. Tronatad and Brun, 1c., see also Collas, Neture, 1933, 142, 569, Bell and Wolfel, and Collas, Neture, 1933, 142, 569, Bell and Wolfel, and Collas, Neture, 1934, 143, 25. Froe Roy. Soc. 1934, 1443, 22. JC.S 1936, 280. Fowler, 1934, 1444, 12. Collaboration, 1934, 1444, 13. Collaboration, 1934, 1444, 13 obtained (cf. Tronstad and Brun, 1 c., see also

(b) Other Methods of Separation of Deutersum. -Deuterium in ordinary bydrogen was first concentrated by utilising the difference in vapour pressure of the liquids. Brickwedde and Murphy (Physical Rev. 1932, 39, 164, 664; 40, 464) used this method and found that the relative abundance of deuterium in the last fraction was increased five or six times compared with the original gas. Keesom, van 1938, 44, 204). The overpotential at a mercury Dijk and Haantjes (J. Proc. Acad. Sci. Amster- electrode in ordinary and beavy water has been dam, 1933, 36, 248) used a specially designed investigated by Heyrovsky (Chem. Listy, 1937, total, 1805, 505, 269) 380 a aprenay seagged investigated by heyrorasy, 1806, 269 aprenay series and 1805, 1805, 269 are also Bowden and Kenyon (Nature, contaming 1-57). Dr. Taylor, Gould and 1835, 185, 195). Bleakney (Physical Rev. 1933, 34, 3498) record in the refractive index, n, for Dg at \$2.662 are not seen to the contamination of the con Amall separation of isotopes by firectional decorption from charcoal. See also Washest and bmith (J. Chem. Physics, 1932, 1, 426), are the Larset (Z. Physik, 1936, 100, 543). Lawson (Trans. Faraday Soc. 1936, 22, 136), are the Larset (Z. Physik, 1936, 100, 543). Lawson (Wat. 1936, 22, 1).

Hertz using his historic apparatus devised pure denterium hy a fractional diffusion method. this vol., p. 605).

The diffusion of deuterium through palladium

was investigated by A. and L. Farkas (Proc. Roy. Soc. 1934, 144A, 467; Nature, 1933, 132, 694); cf. Harris, Jost and Pearse (Proc. Nat. Acad. Sci 1933, 19, 991). Fractionation by diffusion through other metals was examined by Fink, Urey and Lake (J. Chem. Physics, 1934, 2, 105); Jost and Widmann (Z. physikal, Chem. 1935, [B], 29, 247); Luhr and Harris (Physical Rev. 1934, (n), 45, 843); Sieverts and Zapf (Z physikal. Chem. 1935, 174, 559). The fractional distillation of water has been

attempted by Washburn and Smith (J. Chem Physics, 1933, 1, 426); Lewis and Cornish A detailed account of a method using (J Amer Chem Soc. 1933, 55, 2616); Hall and Jones (161d, 1934, 56, 749); Erlenmeyer and

> The rapid conversion of deuterium oxide into deuterium by passing D.O vapour over magnesium at 480°, giving a 95% yield, is described by Knowlton and Rossini (J. Res.

103 ; 200 ; A and L Farsas, 64d 1934; 52 408 ; Urey, Science, 1933, 76, 566 , Eyring, 5 700. Nat. Acad. Sci 1933, 19, 76 ; Eyring and 5 6thers, J. Chem. Physics, 1933, 1, 345; J. 39, 269 ; Physical 2, 1934, 25, 969; Nature, 1934, 2, 217; Nature, 1934, 133, 447; 1938, 44, 21]; Beutler (Z. Physical 1934, 1935, 12, 57, 372); Bul (Nature, 1936, 133, 723); Nothdurft (Ann. Physik, 1937, (v), 28, 157); Megaw and Simon (Nature, 1936, 138, 244), Archer (1814. 136, 286); Grew and Atkins (Proc. Phys. Soc. 1926, 46, 415); Coppock (Trans. Faraday Soc. 1935, 31, 913).

The normal potential of deuterium is given as 44 mv. by Abel and Redlich (Z. Elektrochem.

	D ₂	HD	H ₂
Heat of vaporisation at 194.5 mm. Hg Heat of sublimation at 0°Abs Heat of fusion \theta value for Cp (Debye) \theta value for Cv (Debye) \theta value for Cv (Debye)	47-0 cal. 89° 97° 18-65° 23-6° 128-5 40-5 kg./cm. 23-14 cm. ³ 20-48 cm. ³ ~215 cal.	263 cal. ~228 cal. 37 cal. ————————————————————————————————————	219·7 cal. 183·4 cal. 28·0 cal. 91° 105° 13·95° 20·38° 53·8 30·3 kg./cm.² 26·15 cm.³ 23·31 cm.³ ~305 cal. (5·0±)10-4 0·12

times that of hydrogen. Keyes has calculated η for hydrogen by the method of least squares and finds $\eta = 91.6$ c.g.s. units at 30°, therefore for $D_2 \eta = \sqrt{2} \times 91.6 = 129.5$ e.g.s. units.

Torrey (Physical Rev. 1935, [ii], 47, 644), using Rankine's method, gets $\eta_D: \eta_H = 1.410 \pm 0.03$. See also van Cleave and Maass (Canadian J. Res. 1935, 13, B, 384).

Vapour pressure of HD and D2 was measured by Brickwedde and others (Physical Rev. 1934, [ii], 45, 565; 1935, [ii], 48, 483); Steiner (Z. Physik, 1932, 79, 601); A. and L. Farkas and Harteck (Physikal. Z. 1936, 37, 447); Bartholomé and Eucken (Z. Elektrochem, 1936, 42, 547); the following table is taken from Z. Elektrochem. 1938, 44, 23.

V.P. in mm Hg. H_2 D₂ To Abs. 96.6 12.6 15 16 25.9 155.4 17 237.749.4 18 348.788.2 19 493.7146.3221.0 678-3

For the adsorption of hydrogen isotopes on carbon, copper, lead, nickel and other surfaces, see Barrer (Trans. Faraday Soc. 1936, 32, 481); Beebe and others (J. Amer. Chem. Soc. 1935, 57, 2527); Maxted and Moon (J.C.S. 1936, 1542); Magnus and Sartori (Z. physikal. Chem. 1936, A, **175**, 329); Lennard-Jones and others (Proc. Roy. Soc. 1935, A, 150, 442; 1936, A, 156, 16); Pace and Taylor (J. Chem. Physics, 1934, 2, 578); Klar (Naturwiss. 1934, 22, 822; Z. physikal. Chem. 1935, 174, 1); Kohlschutter (ibid. 1934, 170, 300); Melville and Rideal (Proc. Roy. Soc. 1935, A, 153, 77, 89); Hudson and

Ogden (Nature, 1938, 142, 476).

General Reactivity of Deuterium. Deuterium differs from hydrogen in its chemical reactivity. These differences are sometimes considerable, as is shown by the velocity constant data given in Table A on p. 564. See Geib, Z. Electrochem. 1938, 44, 86, for bibliography and fuller details.

Compounds of Deuterium.-The most completely studied compound of deuterium is the oxide, D2O. Its more important physical constants are given in Table B on p. 564.

For the diamagnetism of D2O, see Gray and Cruickshank (Nature, 1935, 135, 268); Specchia and Dascola (Nuova Cim. 1935, 12, 606).

For dissociation constants of deuterium oxide and of electrolytes dissolved in it, see Schwarzenbach (Z. Elektrochem. 1938, 44, 46); reaction velocity, Reitz (ibid. 1938, 44, 72); thermal properties and dissolving power, Lange (ibid. 1938, **44,** 31).

Reactions in Deuterium Oxide-The substitution of deuterium oxide for water affects the reaction velocity in three principal ways:

I. In the majority of instances so far investigated the effect of heavy water has been to reduce the rate of reaction in solution by as much as 20%. The velocity of a reaction depends to a farreaching extent on the solvent used, even when this takes no active part in the reaction. That there is connection between such physical properties as viscosity, dielectric constant, etc., of the solvent and its influence on the course of a reaction is evident although it has not been possible, so far, to decide as to the quantitative

effect of any particular factor.

2. The interchange between hydrogen atoms of the reactant and deuterium atoms in the water is a further cause of change in reaction velocity. In aqueous solution this interchange is in many instances instantaneous and complete (e.g. $NH_3+D_2O \rightleftharpoons ND_3+H_2O$), with the result that there is a change in the nature of the reacting substances. In other cases interchange takes place in varying extents and rates depending on the experimental conditions, as is notably the case with hydrogen linked to carbon. In general the ultimate effect is a slowing down of the reaction velocity

3. The third effect is produced when deuterium oxide itself takes part in the reaction either as D₂O molecules or in the form of D or OD ions. In this instance the reaction is by no means always retarded; for the most part an acceleration is observed. This is notably the case in acid catalysis and frequently with bases. For a detailed account of reaction velocities in deuterium oxide and bibliography, see Reitz, (Z. Elektrochem. 1938, 44, 72).

Electrolytic dissociation is in general less in D₂O than in H₂O. Typical values are given in Table B, p. 564, but for theoretical treatment of the results obtained together with references

DEUTERIUM.

TABLE A.

						IADLE A.			
		_	 	_				Difference	in kg cal.
Reactions compared Temp. °C.		Temp. °C.	Vel. const., k ₁ /k ₂	Active energy, Q ₁ -Q ₂ .	Zero point energy, E ₁ -b ₂ .				
H+H+(N D+D+(N	1)	•				20	1-4	0	0
H+H,						630-710	1-85 (mean)	0 52	18
D+H,D H+D,D						(wall reaction) ?	~1.7	~0 75	36
$\frac{D+NH}{H+ND}$						260-400	3 2-2	1.5	56
D+PH, H+PD,						420-620	1-04	0.6	43
H+HCI D+DCI						_	-	06	1-2
Na+CIH Na+CID						23B	1-33	0 35	12
CI+HD						30	33	06	0.8
CI+HH CI+DD						0 and 30	13-4 and 0-75	1.2	18
Br+HH Br+DD						275-380	5-5-4-4	1.5	18
$\frac{I_3+H_3}{I_3+D_3}$.						425-500	2 45-2-1	0 75	18
2H1 2DI						430	1.53	0.6	2 093
C ₂ H ₄ +H ₂ C ₂ H ₄ +D ₂						530-570	2.5	0 95	1.8

1,+D,		N.	4	125-	500		1	2	45-2-1	0 75	18
2DI · ·		-1		43	0		1		1-53	0.6	2 003
C ₂ H ₄ +D ₂ · ·		1			570		-		2.5	0 95	1-8
• M is the thir	d body ne	cessar	y for		OCCI			of an	ich a reaction in t	he gas ph	Ase.
			-		ты	-	ь.	_	н,о	T	D,O
Crystalline structure .	٠.		-		•		•		a=4.525 a c=7.39		4-505 A 7-36
Density, d ²⁵ · · ·									0-99705		048
Relative molecular vo	s, at 20°	٠.							1,0037	1	
M.p. B.p. Temperature of max. Heat of sublimation (t Heat of fusion Dielectric const. 0°C. Mol. magnetic suscept:	٠.	٠.				٠			0°C.	3 8	
Dp	lanester	٠.	•		•	٠	٠	•	100°C. 4°C.		-42°C. 5°C.
Heat of sublimation (t	rinle not	nt) .	•	•	•	•	٠		12170 cal./mo		31 cal/mol
Heat of fusion	. Por				•	:	•	٠,	1435 cal./mol.	152	3 cal /mol
Dielectric const. 0°C.					÷			.	81.5	80	7
hiol. magnetic suscept	bility 20	٠.							12 96× 10°		97×10°
Surface tension 12°C. Viscosity 25°C. Refractive index, np		•					٠	٠.	73.7 dynes/cm	. 73	dynes/cm
Viscosity 25 C.		•						٠,	0-891 centipos		92 centipos
		• •	_•		•	•	Ŀ	·	I-33300	1.3	2844
Ionic mobilities, T=18	°C.				-	٠.		7		54.5	CI- 55
Solvent H.D	: :		•	•			15			34 2	CI- 65
Solvent 97% D.D. Solvent H.D Solubility 25°C. g /100	g. solver	ıt .	•	•	**				r	P # P	Ç. 00 .
Solvent D.D Solvent H.D .					N		30		BaC	1, 289	(20°C.)
Solvent H.D .					N	C	35	9	BaC	1, 357	(20°C.)
Electrolytic dissociation	תו						_	_		-	
D.D Kp.o=016x	10-11	٠.		٠	K	40	-1	6x	10-11		
HDD (D+)=0-111)	(10.4	٠.			(H	I+]	-0	004	×10-7		

Dissocn. Const. in H2O

trochem. 1938, 44, 46).

Dissocn. Const. in D₂O at 20° C. Substance. 2 HSO₄ н-соон . 2.50 CH_s·COOH 2.87 Hydroquinone . 3.35

Salts having water of crystallisation can have one or more of the molecules of water replaced by a corresponding number of molecules of D2O. Research on the deuterates of CuSO₄ has been carried out by Partington and Stratton (Nature, 1936, 137, 1075); Mules and Menzies (J. Amer. Chem. Soc. 1938, 60, 87), and by Perpérot and Schacherl (J. Phys. Radium, 1935, 6, 439). Taylor (J. Amer. Chem. Soc. 1934, 46, 2634) transition temperature Na₂SO₄,10D₂O to be 34·48°±0·02° as against 32·38°±0·001° for the corresponding hydrate. Bell (J.C.S. 1937, 459) measured the dissociation pressure of a number of denterates, e.g. CuSO₄,5D₂O; SrCl₂,6D₂O, etc. Godehot, Cauquil and Calas (Compt. rend. 1936, 202, 759) obtained the deuterates of krypton and xenon with 6D2O.

Other Inorganic Compounds of Deuterium.—These can for the most part be prepared by several general methods slightly modified to suit individual cases.

1. Acids, by action of D₂O on the acid anhydride, e.g. D_2SO_4 (Ingold and others, J.C.S.

1936, 916).

2. Non-Metallic Deuterides, from the elements with or without a suitable catalyst, e.g. DBr or by the action of D_2O on a suitable salt, e.g. D_2S from $Al_2S_3+D_2O$; ND_3 from Mg_3N_2+ D₂O.
3. Interchange reaction with or without a

4. Action of D₂O on the metal yields denteroxides in the case of some metals, e.g. NaOD.

For a detailed account of inorganic deuterium compounds and a bibliography, see Erlenmeyer (Z. Élektrochem. 1938, 44, 8).

Some typical differences in the physical preperties of corresponding denterium and hydrogen compounds are shown in the following

rante:				
Com- pound. HCl	M.p. absolute	B.p. absolute	Latent heat of evapn. cal./mol.	Critical temp.
	$162 \cdot 2$	188-1	4081	51.0
DCI	158.2	191.6	4151 ′	50.3
HBr	186.2	206.3	4257	89-9
DBr	185.7	206·3	4258	88.8
HI Di	222·3 221·5	237·5 237·0	4724 ? 4713 ?	150·7 148·6
	221.0	237.0	41191	140.0
NH_3	$195 \cdot 3$	239.8	5797	132.5
DH ₃	199.6	242-1	5990	132.3
HCN	259	298.5	6600	
DCN	261	299.1	6500	

Organic Compounds of Deuterium.—The may be carried out by a large variety of pre- deuterium occur very rapidly, in other cases the

to the literature, see Schwarzenbach (Z. Elek- | parative methods, many of which are exactly parallel to the general methods of preparation of the corresponding hydrogen derivatives.

Examples of these are:

1. Treatment of carbides with D2O.

2. Catalytic deuterisation.

3. Grignard reactions.

4. Addition of D₂ under suitable conditions.
5. Exchange reactions.

The following are examples illustrative of some of these methods:

Tetradeuteromethane, CD4, is prepared by the reaction of D₂O with aluminium carbide (Urey and Price, J. Chem. Physics, 1934, 2, 300).

Mono deutero-tetramethylmethane,

(CH₂)₂C·CH₂D,

was prepared from D₂O and the Grignard compound of 1-chlor-2:2-dimethylpropane, (Whitemore, Fleming, Rank, Bordner and Larsén, J. Amer. Chem. Soc. 1934, 56, 749, 934). For a comprehensive bibliography up to January, 1938, see Erlenmeyer (Z. Elektrochem. 1938, 44, 9).

Differences in melting-point and boiling-point of corresponding deutero- and hydrogen organic compounds are shown in the following table:

Substance.	M.p. °C.	В.р. °С.
C ₆ D ₆	5·5 6·8	80·1 79·4
C ₁₀ H ₈ C ₁₀ D ₈	80·2 77·5	
CH ₃ ·CHO CD ₃ ·CDO (CH ₃ ·CHO) ₃ (CD ₃ ·CDO) ₃	-123·5 -121·7 10·5 13·7	20·2 (760) 20·5 (756) 124 (760) 124 (753)
CH ₃ ·COOH CH ₃ ·COOD CD ₃ ·COOD CD ₃ ·COOD	16·6 15·4 17·2 15·9	
CH₄ CD₄	-90·6 -89·2	

Interchange Reactions.-These may be divided into two classes, homogeneous and heterogeneous.

Homogeneous Reactions .- A number of substances containing hydrogen exchange one or more of their hydrogen atoms for deuterium when dissolved in D₂O. Certain of these arc obviously ionic reactions and, as such, occur very rapidly, e.g. on dissolving sodium hydroxide in deuterium oxide interchange occurs instantaneously according to the equation:

NaOH+D₂O ⇒ NaOD+HDO

This exchange can be followed quantitatively by recovering the diluted D_2O subsequently and determining its density. In a mixture of H_2O and D_2O the available deuterium distributes itself between solvent and solute in proportion to the molecular quantities of each present. preparation of organic compounds of deuterium For the most part the exchange reactions of exchange takes place with messurable velocity, fractionation of H2O and D2O in the organism and an analysis of the kinetics of such reactions shows that the ionic reaction of hydrogen interchanga itself must be preceded by some intermediate reaction the velocity of which determines the rate of the whole reaction. Amongst slow morganic interchange reactions the exchange of hydrogen in complex salts should be noted. Hexamminecobaltic chloride, for instance, inter changes its hydrogen for deuterium and it has been found that the reaction velocity is inversely proportional to the hydrogen ion concentration. This discovery is of importance in atudying the theory of complex salta (see Erlenmeyer and Gärtner, Helv. Chim. Acta, 1934, 17, 1008; Bankowski, Monatsh. 1935, 65, 266, James, Anderson and Briscoe, Nature, 1937, 139, 109; Anderson, Spoor and Briscoe, shid 1937, 139, 508; Garrick, sbid, 1937, 139, 507).

Bonhoeffer and Brown (Z. physikal Chem. 1933, B, 23, 172) were the first to discover the interchange phenomenon. They found that NH₄Cl dissolved in D₂D exchanged all its hydrogen atoms, and this reaction has been utilised to ohtain water free from denterium It has been found that hydrogen which is linked to oxygen, nitrogen, sulphur or halogen readily interchanges with deuterium. Since then it has been determined that many other aubstances, e g. glycol, hydrogen peroxide, acetoacetic ester, phenols, etc., exchange one or more hydrogen atoms more or less readily according to the conditions of experiment. For hibbography, see Ingold and Wilson (Z Elektrochem, 1938, 44,

A limited number of interchange reactions occur with deuterium atoms in the gaseous phase. The source of such atoms may be the electric discharge, photochemical excitation by means of mercury resonance radiation, or thermal dissociation of heavy water molecules with mai discolation of neary water molecules with a land syntogen in the painter aeros are in agree formation of atomic deuterium (Gels and Straces, Z., physikal, Chem. 1935, B. 29, 216; b. Staceic and Phillips, J. Chem. Physics, 1936, 1834 and on the Rydberg formula. Staceics, J. Tenneer, Taylor and others, stud. 1937, 1832, 892; J.C.S. 1936, 26; Proc. Roy. Based on the thooly of the isotope effect. The molecules and phytogen accord with electhations 1933, 1828, 892; J.C.S. 1936, 26; Proc. Roy. Based on the theory of the isotope effect. The control of the control of the second with electhation of the control of the c

Soc. 1936, 157, A, 625).

Heterogeneous Interchange.—Interchange of Heterogeneous increases ample molecules is catalysed by the common hydrogenation catalysts such as platinum, palladium, nickel, iron, etc. (Farkas, and Farkas and others, Trans Faraday Soc. 1935, 31, 821; 1936, 32, 416, 922; J. Amer. Chem. Soc. 1938, 60, 22; Taylor and others, 15id. 1935, 57, 680, 1256, 1936, 58, 1445; 1938, 60, 362; Hirots and Horinti, Sci. Papers Inst. Phys. Chem. Res. Tokyo, 1936, 30, 151; Horiuti and Polanyi, Trans Faraday Soc. 1934, 80, 1164; see also Ingold and Wilson, Z. Elektrochem, 1938, 44, 62)

Blological Applications. - The use of denterium as an indicator in the study of intermediary metabolism has produced much useful

could be detected in normal cases. Woglom and Weber (J. Amer. Chem. Soc. 1934, 104, 1289) showed that D.D has no effect on mouse sarcoms. Deuterium has been used as an indicator in fat and other metabolism (Cavanagh and Raper, Nature, 1936, 137, 233; Rittenberg and others, J. Biol. Chem. 1936, 114, 381; 115, 635; 1937, 117, 485).

The incorporation of deuterium in the hving organism is the subject of a research by Bon.

hoeffer and others (Z. physikal, Chem. 1936, 175, 459; 176, 202; 1937, 180, 185). The effect of exchange of deuterium for hydrogen in experiments involving enzymes and moulds is to produce a greater or less reduction in activity. Pollen germinates more slowly in O₂D than in ordinary water (Plantefol and Champetier, Compt. rend. 1935, 200, 423)
Yeast is less active in presence of deutenum orade (Taylor and Harvey, Proc. Soc. Exp. Bol. Med. 1934, 31, 954; cp. Hughes, Yndkin, Kemp and Rideal, J.C.S. 1934, 1105; Steece, Z. physikal, Chem. 1934, B. 27, 6; Bonhoeffer and Salzer, Naturwiss, 1935, 23, 867; Shoup and Meyer, J Tennessee Acad. Sci. 1935, 10, 127; Salzer and Bonhoeffer, Z. physikal Chem. 1936, 175, 304; von Dungern, Z. Biol. 1936, 97,

A summary of researches in physiological ehemistry using deuterium compounds is given by Bonhoeffer (Z. Elektrochem, 1938, 44, 87)

and Theis (Woch, Brau, 1938, 55, 36). Spectroscopic Data,-The wave lengths of chief lines in the Balmer series of deuterium are given below, the bracketed data being values for the corresponding Ha, Hg, Hy, and H& hnes Da 6561 00 (6562 793); Dg, 4880 00 (4861-326); Dy, 4339 282 (4340 487), and D3 4100 619 (4101 738). The differences between deuterlum and hydrogen in the Balmer aeries are in agree-

spectra of the numerous other distomic molecules have also been examined (e.g. LID, NaD, AgD, CaD, AiD, etc.). For data, see "Tables Annuelles de Constantes et Données Numériques, l Deuterium et Composés de Deuterium," by

G. Champetier (Hermann & Cie, Paris), 1937.
The general effect of substitution of deuterium for hydrogen as to change the features of the spectrum dependent on the moment of inertia of the molecule The interatomic distance is not altered by substitution of D for H. The change in the moment of inertia may have profound effect on the general appearance of the spectrum Thus the ultra violet absorption spectra of the molecules ND, and NH, or of CH, ND, and CH, NH, are quite distinct, sa ara tha infra-red apectra of molecules such as DCl and HCl information. von Hevesy and Hofer (Nature, or D.) and H.O. These distinctions have also 1934, 1934, 879) used deuterium oxids to follow been observed in the Raman spectra of a large manner of authances, C., H.D. and C.H., or Elemanyer and Gariner (bibl. 1006) examined its C.H., C.D.D. C.H., C.D.H. and C.D., C.O.D. distribution in rats. Breusch and Hofer (Berlin. For comprehensive data, see Champetier, op cit.) klin. Woch. 1934, 13, 1815) found that no The table below illustrates typical changes in brought ahout by substitution of D for H.

folecule.	Fundamental frequencies, cm.—1							
H ₂ S .				. 2578				
D ₂ S .				. 1875				
H ₃ As.				. 2094				
•				990				
				910				
D_3As .	-			. 1508				
-				730				
				630				

In the case of more complex molecules (e.g. with C6H6 or CH3·COOH) partial suhstitution of hydrogen hy deuterium has been used as a means of assigning observed vibrational frequencies to the appropriate molecular vibration.

Estimation of Deuterium.—A number of methods are in use for the estimation of D. some of which can be applied only to water and

others only to gaseous hydrogen.

Spectroscopic and Mass Spectrographic Methods. —These methods are qualitative as well as quantitative. The first quantitative estimation of deuterium was carried out spectroscopically by Urey, Brickwedde and Murphy (Physical Rev. 1932, 39, 164, 864). The method was based on the different wave-lengths of the corresponding lines in the atomic spectra of the hydrogen isotopes and was carried out by comparison of the intensities of the β and γ lines of the Balmer spectrum. Bleakney (ibid. 1932, 40, 496; 41, 32; 1933, 44, 265; 1934, 45, 281, 655) devised a very accurate method using the mass spectrograph and which required only small quantities of gas.

A thermo-conductivity-micro method employed by Farkas for which 2-3 cu. mm. of gas were required. The D content, however, must not be less than 1%. It has an accuracy of ±0.1% (see Farkas, Proc. Roy. Soc. 1934, 144, A, 467; Trans. Faraday Soc. 1936, 32, 413; Farkas, Farkas and Rideal, Nature, 1936, 137, 315; see also Harteck, Z. Elektrochem. 1938,

The most widely used method of estimating deuterium is that depending on the direct determination of the density of water containing deuterium either by the pyknometer or hy means of the float, each of which is capable of an accuracy of 1 in 10⁻⁶. This method is applicable to the determination of D₂ in all compounds which can be oxidised to give water. For details of the pyknometric method, see Washhurn and others (J. Res. Bur. Stand. 1933, 11, 453; 1934, 12, 305). For the float method, see Lewis and MacDonald (J. Chem. Phys. 1933, 1, 341); Emeléus, James, King, Pearson, Purcell and Briscoe (J.C.S. 1934, 1207; 1935, 1545). Bibliography.—A. Farkas, "Orthohydrogen, Parahydrogen, and Heavy Hydrogen," Univer-

sity Press, Cambridge, 1935; G. Champetier, "Tables Annuelles de Constantes et Données Numériques. I. Deuterium et Composés de Deuterium," Paris, 1937.

the fundamental frequencies of molecules 15% of zinc. Devarda's alloy reduces nitrates to ammonia quantitatively in alkaline solution, the ammonia being then distilled into standard acid solution (Devarda, Chem.-Ztg. 1892, 16, 1952; Z. anal. Chem. 1894, 33, 113; Busvold, Chem.-Ztg. 1914, 38, 799) (v. CHEMICAL ANALYSIS, Vol. II, pp. 598, 664).

DEXTRAN, C₆H₁₀O₅, or viscose, is a gum

which occurs in the unripe sugar beet (Scheihler, Wag. J. 1875, 790). It is formed in the lactic fermentation of cane sugar hy the action of Streptococcus (Leuconostoc) mesenteroïdes, (van Tieghem, Jahresber. Agric. Chem. 1879, 544; Bechamp, Compt. rend. 1881, 93, 78; Brüning, Annalen, 1857, 104, 197). S. hornensis isolated from curdled milk, some water samples and flowers also converts media containing up to 20% sucrose into dextran (Boekhout, Centr. Bakt. 1900, [ii], 6, 161).

An animal dextran, $C_6H_{10}O_5$, is found in the galls produced on elms by the louse (Schizoneura lanuginosa) (Liebermann, Pflüger's Archiv. 40, 454).

DEXTRINS, $n(C_6H_{10}O_5)$. When diastase acts on starch paste, maltose is the principal product, hut various dextrins are also formed, their nature depending on the nature of the enzyme and the conditions of action. The precise nature of these products of the degradation of starch still remains obscure and it is not easy to make a satisfactory review of the subject at the present time. Recent work, however, points strongly in the direction of the dextrins being side products or by-products of the action of enzymes on starch rather than, as previously believed, intermediate products formed on the way to maltose. The dextrins differ according to the conditions ($p_{\rm B}$, temperature, concentration) and the nature of the enzyme used. Some (like a-amylodextrin) appear to be fragments of the original starch moleculo left unattacked by the enzyme preparation. Others appear to be formed owing to reversion or retrogradation occurring (stable dextrins).

According to their behaviour towards iodine

they have been classed into:1

Amylodextrins, giving a blue colour and soluble in 25% alcohol.

Erythrodextrins, giving a reddish-brown colour and soluble in 55% alcohol.

Achroodextrins, showing no colour and soluble

in 70% alcohol.

In the following a summary is given of the more definite dextrins which have significance in regard to the structure of starch and the mode of action of the two amylases (the α - and β -

amylases of recent workers).

Unfortunately Syniewski, whose work perhaps forms one of the best contributions to the subject, has used a different and special nomenclature. Especially interesting in Syniowski's work is the proof that ungerminated barley is actually richer in diastaso (his a-diastase) than the malt made from it. This diastase, however, is present in a form which is insoluble in water and is only hrought out hy digestion with a

DEVARDA'S ALLOY. A hrittle alloy containing 45% of aluminium, 50% of copper, and 1 An account of starch-iodine coloration as an index of differential degradation by the amylases has been given by C. S. Hanes and M. Cattle. Proc. Roy. Soc. 1933, B, 125, 387-414.

protease—papayotin; during germination the It is converted by malt extract previously heated barley amylase is apparently released and partly lat 76°C. into 67°3% maltoss and the reducing converted into the second amylase (his \$d_{obs} | Lumt-detritin 1 (ca]=180°). tase). He shows how the two diastases augment each others' activity, and works out a quantitative method for measuring this effect.

The dextrans are characterised physically by their optical rotation and cupric reducing

power.

The early work of Brown, O'Sulhyan, etc. gave the reducing values (K) in terms of glucose taken as 100. Later values were expressed in terms of maltose as 100 by the symbol R, or preferably RM.

In Cermany and the USA, the West method and tables (" Tabellen zur quantitativen Bestimmung der Zuckerarten," 1888) are still used m estimating maltose in spite of the fact that British workers since 1897 have chown them to be incorrect to the extent of being 5% low for anhydrous maltose (Brown et al., J.CS. 1897, 71, 103-196; Ling and Baker, thid 509)

Symewski since 1925 has used corrected Wein

I. THE SO CALLED AMYLODEXTRING (PRODUCING A PUBE BLUE COLOUR WITH IODINE) (NON-

A .- The so-called "amylodextrin" or solubilised starch (Symewski, Annalen, 1902, 324, 212; 1925, 441, 277), is made from potato starch by hesting with water in an autoclave (12 hours at 140°C.) until it is completely soluble.

It has the composition CateHand iss, and was regarded by Symewski as 12 maltone radicals

combined with a destructing complex.

B—a. Amylodextrin, [a]p 190-195°, Rg=
0 50-2 (J. L. Bsker, J C S 1902, 8t, 1177), is prepared by the action of precipitated diastase from barley extrect (i.e. the camplase of Syniewski, Biochem. Z 1925, 158, 87) on Lintner's soluble starch, or on starch paste, pre ferably at 50° C. (The action on starch paste is slower than that on soluble etarch) It is sparingly soluble in cold water, readily in hot water. Solutions give a pure blue colour with todine. It is converted by precipitated malt diastase efter 18 hours' action into maltose. achroodextrm, and a considerable amount of glucose. C. Non-reducing Limit-dextrin i (Grenz-

dextrin 1), [a]n+194 92°, R =0 97 (Symewski, Annalen, 1925, 441, 285; he was unaware of J. L. Baker's work). Prepared by the action of barley extract (i e. the a amylase of Symewski, Biochem, Z. 1925, 158, 87) on his amylodextrin solutions (5 7%) at ordinary temperature in 75 minutes; action then became very slow. The alight reducing power is regarded as due

to decomposition of the dextrin by hot Fehling's solution. It gives a pure blue colour with iodine solution

Analysia indicated:

It polymerises in cold aqueous solution to all the molecular weight about 52 (G₂, H₁₁, G₂), and the molecular weight about 52, bow The value given mixed (wield by the Y.P. method gives 6,002 (Theory 5,991).

The a-amylodextrin of Baker or non-reducing Lamst-dextrus I of Syniewski would be the a-

atarch of Van Klinkenberg (Erg. Enzymforsch. 1934, 3, 71) left unattacked by his \$ diaptase. The above aubstances must be distinguished

from Nageli's amylodextrin (1874), which is obtained by the very prolonged action of scids on potato etarch at the ordinary temperature This, according to Brown and Morns (JCS. 1889, 55, 449) is easily coluble in hot water, dessolves in cold water to form a 1.75% solution. and produces an intense-red coloration with nodme. It had [a], zes=206 3°, Kzes=907, and was completely converted into maltose by maît extract in 20 minutes, at temperatures up to 60°C This dextrin was a strongly reducing cubstance,

Lintner and Dull'a "amylodextrin" (Ber. 1893, 26, 2533) or "amorphous soluble starch" had feln=196° and was non reducing and apparently the same as Baker's and Syniewski's materral.

II. REDUCING DEXTRINS-"LIMIT DEX-TRINS " (GRENZ DEXTRIN).

D—Limit-dextrin1, $[a]_D^{20}+1796^\circ$, $R_N=1765$ (Wein) 1 and 16 2° (Wein) (Syniewski, Annalen, 1899, 309, 282; 1902, 324, 212), is prepared by the action of fresh malt extract on starch paste at ordinary temperatures, and sisc on solutions of amylodextrins.

Analysis indicates .

Mol. wt. found=1,897 (F.P.). (Theory=1,980) When treated with mait extract for a very long period (three weeks in 5% solution) it gave "isomaltose," which was axtremely bygroscopic, [a] =+141-4°, Rx=845 (Wein), osazone, m p. 152 5°.

Symewaki terms this isomaltose " dextrinose." E-Limit-dextrin 11,[a]p+179 6°, Rx=30 0 (Wein) (Symiewski, Annalen, 1902, 324, 212). is made by the action of malt extract previously heated for 15 minutes at 78-77°C this would be by the action of the \$ disstass of Synicwski, the a diastase of others) on solubilised starch, until the rodine colour just disappears Analysis indicated:

$C_{34}H_{62}O_{31}=8(C_6H_{10}O_5),H_2O_6$

Mol. wt. found = 1,039. (Theory = 990) It dessolves fairly readily in hot 80% alcohol, aparingly in 90% alcohol, and is converted slowly by unheated fresh malt extract (45°C., 48 hours) though nearly quantitatively to maltose (990%) Wein=101% corrected).

This is identical with the following dextrins of other workers:

E1-Maltodextrin, [a]; 2 m=193.1°, Kin =21-1 (Brown and Morris, J.C.S. 1885, 47, 527).

DEXTRINS.

made by the action of freshly prepared diastase from green malt on starch paste at 60-65°C. (This would be the action of the β -diastase of Syniewski, the action being carried to [a], 3.86= 198°.)

Brown and Millar (ibid. 1899, 75, 286) give the equivalent $[a]_D=180^\circ$, $R_M=34\cdot5$. It was regarded as $C_{12}H_{22}O_{11}<(C_{12}H_{20}O_{10})_2$ and was said to be completely convertible by fresh diastase into maltose.

 E_2 —Maltodextrin-a, [a]_D 180°, R_M =32·8 (Ling and Baker, *ibid*. 1897, 71, 508), was made by the action of precipitated malt diastase on starch paste at 70°C., and had the formula C₃₆H₆₂O₃₁, mol. wt. found 990.

It is sparingly soluble in 80% alcohol, and is said to be incompletely converted to maltose by precipitated malt diastase. (After 48 hours at

room temperature the conversion had [a]p 3.93= 141·3°, R_{3·93}=90·8.)

 E_3 .—Achroodextrin II, $[a]_D$ 183°, $R_M=$ 26.5-26.8 (Lintner and Düll, Z. gcs. Brauw. 1894, 17, 339), was separated from starch conversions. They regard Brown and Morris's maltodextrin as a mixture of achroodextrin II and "isomaltose."

III. SIMPLER DEXTRINS.

F.—y-Maltodextrin, $[a]_D$ 172.28° solution), R_M=42.7 (Wein) (Syniewski, Annalen, 1902, 324, 212), made by fresh malt extract acting on Limit-dextrin II for one hour until the product shows 60% apparent maltose and then fractionated. It is easily soluble in 80% alcohol, sparingly in 90% alcohol. Analysis indicated $C_{24}H_{42}O_{21}=4(C_6H_{10}O_5)H_2O$. Mol. wt. found =595 (theory 666). It is said to give maltose and some "isomaltose" when saccharified by fresh malt extract at 45°C.

It is identical with the following:

 F_1 .—Maltodextrin-β, [α]_D 171·6°, R_M = 43·0 (Ling and Baker, J.C.S. 1897, 71, 508), obtained by starch transformations using precipitated malt diastase at 70°C. (Action of Syniewski's β -diastase, Kuln's α -diastase.) Formula: $C_{24}H_{42}O_{21}$. Mol. wt. found=670. It is incompletely converted by precipitated malt diastase in 24 hours. Product had $[a]_D=140^\circ$, $R_{M}=91.5$.

F₂.—Achroodextrin III, [a]_D 171·1°, R_M= 42·5 (Prior and Wiegmann, Z. angew. Chem. 1900, 13, 464) which is made by the incomplete conversion of potato starch paste by an extract of green malt (giving a red colour with iodine). Formula: $(C_{12}H_{22}O_{11})_2, H_2O$. It differs from Lintner and Düll's achroodextrin II in being partly fermentable by Saaz yeast and more fermentable by Frohberg yeast, by which in vacuo it is said to be completely fermentable. It is completely converted by Logos yeast. With diastase (½ hour at 50°, gradually rising to 60° and 70°C.) it gave maltose and Prior's achroodextrin IV, C₁₂H₂₀O₁₀, H₂O, but no "isomaltose." F₃.—"Maltodextrin," [a]_D 181·3°, R_M=42·5 (Brown and Millar, J.C.S. 1899, 75, 286), which

was made by the action of kiln-dried malt (D.P. 38°L.) i.e. by the action of Syniewski's β -diastase, Kuhn's α -diastase on highly concen-

trated starch solutions (15%) at 55°C.

This has the same R_M as Ling and Baker's maltodextrin- β , but higher $[a]_{p}$. It is said to give mallose only with fresh diastase, without the formation of "isomaltose" or stable dextrin. It is not in the slightest degree fermentable by (English) brewery yeast.

IV. THE SO-CALLED "STABLE DEXTRINS."

G.—Stable-dextrin, $[a]_D$ 197–198°, $R_M=5.5$ (Brown and Millar, J.C.S. 1899, 75, 315), is made by the action of cold water extract of malt on gelatinised starch at temperatures below 60°C, the action being carried to the "resting stage" ($[a]_D=150^\circ$, $R_M=80$).

It is not resolvable by fractionation. Unlike the maltodextrins, it is attacked very slowly by large quantities of malt extract. A 7% solution was only saccharified to the extent of 30% after 48 hours by 25 c.c. of malt extract per 100 c.c. of solution. (The malt extract was prepared from a kiln-dried malt and found to be free from maltase.) The product consisted of equal parts of maltose and glucose. It was regarded as 39 C₆H₁₀O₅ groups and IC₆H₁₂O₆ (aldehydic). Mol. wt.=6,221.

H.—Stable-dextrin, $[a]_D$ 185°, R_M =14 (Ling and Nanji, J.C.S. 1925, 127, 636), which is made by the saccharification of potato starch by precipitated malt diastase (from brewer's malt Z.V. 30°) at 40°C. until the apparent maltose was 80.8%. The product was carefully frac-tionated. M.wt.=1,923. It is regarded as a tetra-amylose and said to be transformed slowly by maltase into isomaltose and with emulsin to give a mixture of maltose and glucose.

It is clearly quite a different material from

Brown and Millar's stable dextrin.

Schardinger by the action of macerans on starch obtained crystalline dextrins designated as a- and \(\beta\)- which Pringsheim has termed polyamyloses.

Pringsheim sets out the polyamyloses as

a-series. Rotation. a-hexa-amylose $[(C_6H_{10}O_5)_2]_3 + 139^\circ$ a-tetra-amylose $[(C_6H_{10}O_5)_2]_2 + 148.5^\circ$ di-amylose $(C_6H_{10}O_5)_2$ +136.5° β-series.

 β -hexa-amylose $[(C_6H_{10}O_5)_3]_2 + 158^\circ$ Tri-amylose $(C_6H_{10}O_5)_3$

and describes their esters and halogen addition products.

Karrer and Bürklin (Helv. Chim. Acta, 1922, 5, 181) question the existence of triamylose, and considers it and β -hexa-amylose to be a polymeride of maltose anhydride, i.e. an octaamylose.

These polyamyloses yield glucose on complete hydrolysis, acetylbromide converts them into acetobromomattose, whilst hydrochloric acid and silver carbonate cause the formation

of amylobiose (isomaltose).

.When tetra-amylose is dissolved in formamide and precipitated with alcohol, 80% of crystalline a-amylosan is obtained, and the balance of 20% is obtained as crystalline β -amylosan on distillation of the filtrate (Pringsheim, Wieder and Weidinger, Ber. 1930, 63, [B], 2628).

The composition of these dextrins is as yet] too uncertain to make further discussion of valus. The auhiert has been discussed by Pringaheim ("Die Pelysaccharide," Berlin, 1931; alse "Chemistry of Monosaccharides and Polysacchandes," 1932, pp. 280-295), and hy Pictet (Xème Conférence da l'Union Internationale de Chimia Liège, 1930).

The very properties which make the dextrins obscure chemically, namely, their varied properties, have been turned to advantage in andustry, where they serve a wide variety of

purposes as adhesives.

In industry the processes for the production of dextran may be divided into two classes. (1) Dry starch is subjected to the action of

heat alone or with chemical agents. (2) Wet starch, suspended in water or

cooked to a jelly, is acted on with chemical agents with or without heat

The former is the more important The products vary greatly from almost pure white the product just cases to give a hiue foding to dark hrewn, and there is no close definition coloration, and the products of the further of dextrin, though there is a U.S. Government heating are soluble in cold water. When care is specification. A summary of the subject with taken to exclude moistura, and the heating is an abstract of the literature is given by Bloede | continued for 2 or 3 hours only, using equal

Any one of the aterches in commerce can be used as raw maternal Potato starch is the sasieat to convert and produces the highest type of dextrin, but the taste developed makes it undesirable for postage stamps and envelope gum. Tapieca atarch la preferable, producing raw material for high grade detrim. Corn (maize) starches are inferior for many purposes,

but as they are cheap they exceed all others in

annual tonnage used. The starch is first freed as much as possible frem combined water by drying in suitable ovens, and then anhinited to a temperature of 212-275° in rotating sheet-iron drums. The heat is supplied by hot oil, rape oil being fairly generally used, or by direct fire. When the oil-bath is used, the centrol over the temperature is greatest, and the colour of the product can be varied at will from white to dark brown. When tha drums are heated by direct fire, they are greater viscosity and producing a much thicker made to rotate on slightly inclined axes, the liquid when made up with the same quantity of dried starch being fed in at the higher end of the revolving cylinder and the product discharged at the ether. Sheet iron ovens are also used; they are constructed singly or in sets. The heat is supplied by a furnace, the heated air from which is drawn over the top of the ovens. The material is kept continually stirred by flat iron oars. Boxes and trays are also used as converters, and, indeed, tha maker has only to consider the variety and quality of product required, and to remember that the drier the atarch and the lewer tha temperature of conversion, the whiter will be the product, and, further, that different starches yield different products, i.e. behave differently when suh mitted to the action of heat, to enable him to construct an apparatus that will suit his requirements. It must also be borne in mind that the conversion takes place more slowly at a low temperature,

is required to yield a satisfactory product, is to spray or moisten the atarch either with a mixture of dilute nitric and hydrochloric acids, or with each and singly, or with dilute oxalic acid. The paste is dired and heated at 100-120° or to 150°, till the transformation is complete. as determined by the iodina test; it must then be arrested promptly.

The alternativa method, in which less heat

For a method of examination of commercial dextran and related starch products, v. Babington. Tingle and Watson, J.S.C.I. 1918, 37, 257F. The manufacture of potato dextrin is described by A. E. Williams (Chem, Trada J.

1932, 90, 99; Ind. Chem. 1932, 8, 114). On heating dry atarch with formic or acetic acids, the starch nucleus is broken down and esters of dextrins are formed, which are soluble m cold water. The proportion of acid fixed depends on the time of treatment, After I5 hours' heating with glacial acctic acid at 90°, in Walton's "Comprehensive Survey of Starch weights of starch and glocal accide acid, sn Chemistry," New York, 1923 acetylated starch, known commercially as Feculose, is obtained. This behaves like starch towards boshing water, but the jally does not revert on standing It forms very clear flexible films, and is said to give better results than dextran as a textile finish (see Traquair, J S C I. 1909, 28, 288)

During hydrolysis the formation of glucom should be limited as far as possible, as it increases the hygrescopicity of the dextrin. The quantity present depends to some extant on the natura of the hydrolytic agent, of which 1-30 of 1% is

employed.

The operation of dextrinisation is and to be still largely an art, there being a variable resistance of the starch to the action of the hydrolyang agent, doubtless due to the presence of small hut varying quantities of impurity in the starch. This view is ampported by the fact that the resistance of the lower grades is more marked.

British gum differs from daxtrin in having

water. This is due to the presence of only partly converted starch. For its manufacture rosating alone is generally sufficient, rapid heating to between 350° and 400°F, being

"Gum powders" have recently become of emportance as hottle adhesives and are special

powdered forms of dextrin

Paste or only partly converted dextrins are advantageously made by means of enzymes. The thin bedding starches required by the textile industry are usually prepared by making a cream of the raw starch in water, adding the requisite amount of mineral acid, and digesting at 100-120°F., a temperature below the bursting point of the granules. A better, though more expensive, method is by treating the starch in the cold with an oxidising agent auch as sodium hypochlorite.

Vegetable glue is prepared by alkalina treat-

ment of starch, for example, by heating with | both) are terminally situated and which conse-5% caustic soda with constant agitation to the bursting point of the starch granule. The stringy glue-like mass has high adhesive properties and is widely used in the wood-working industry as a substitute for animal glue.

"DEVIL'S" DUNG v. ASAFŒTIDA.

DEXTROFORM. A non-poisonous combination of dextrin with formaldehyde which is used as a disinfectant dusting powder in the treatment of wounds and infected ulcers.

DEXTROSE, glucose, v. CARBOHYDRATES. DHAK GUM, a variety of Indian kino obtained from Butea frondosa Roxb. (q.v.).

DHURRIN is a cyanophoric glycoside iso-lated by Dunstan and Henry (Phil. Trans. 1902, 199A, 399) from the leaves and stems of the great millet. It is para-hydroxymandelonitrile glucoside and is hydrolysed by emulsin.

E. F. A. DIABASE. This name was introduced by the French mineralogist A. Brongniart, in 1807, for those varieties of basalt in which the light and dark minerals, respectively felspar and augite (or as he erroneously supposed, felspar and hornblende), are distinguishable to the unaided eye. As actually defined by him, the term is therefore synonymous with the diorite of Hauy (1822), whilst with the correction of augite for hornblende it becomes a synonym of the dolerite of Hauy (1822). In the latter sense it is used by some authors at the present time. Unfortunately, the term was re-defined by German petrographers, and it came to be applied to the older, or pre-Tertiary, dolerites, which differ from those of more recent date only in showing correspondingly more alteration, with the development of secondary minerals, such as chlorite. In consequence of this, the name "diabase" is now applied by most English authors to a partially altered or weathered dolerite. In this sense it has an economic bearing, since rocks of this class, being of wide distribution, are much quarried for road making. For this purpose the fresher dolerites are somewhat brittle, whilst diabases are tougher and more durable, providing that decomposi-tion has not proceeded too far. There is also a tendency to use the name "diabase" in a loose sense, like the popular terms "greenstone," "whinstone," "toadstone," and "trap-rock."

DIABETIN. Trade name for a preparation containing 12% of saccharine, 45% of sodium bicarbonate, 43% of tartaric acid and traces of red colouring material (J. Polak, Pharm. Weekblad, 1909, 46, 519).

DIACETIN v. ACETIN.

DIACETYL, CH3·CO·CO·CH3, the first member of the aliphatic 1:2-diketones, a yellow liquid, b.p. 87-88°. The characteristic odour of butter is due to the presence of a small quantity

of diacetyl (about 0.0005%) (v. BUTTER).
DIACETYLENES. The members of this group of hydrocarbons have two acetylenic linkages in the molecule. These linkages may be conjugated or unconjugated. Two principal classes of diacetylenes are to be recognised, In general, unsymmetrical diacetylenic hydroviz. those in which the acctylenic linkages (one or

quently retain one or two reactive ("acetylenio") hydrogen atoms, and those in which all the reactive hydrogen atoms have been substituted by other groups. Substances of the former class (e.g. diacetylene) can display both substitutive and additive reactivity, whereas those of the latter class (e.g. dimethyldiacetylene) can only display additive reactivity.

Preparation.—Diacetylene, CH:C·C:CH, is conveniently prepared by the action of cupric chloride solution on cuprous acetylide, followed by treatment of the copper diacetylide thus formed with dilute hydrochloric acid. The gaseous diacetylene so produced is purified in the Stock vacuum apparatus. Increase in the relative proportion of the cupric chloride, up to a point, increases the purity of the diacetylene produced, but also diminishes the total yield of gas owing to increased oxidation of the copper compounds (Straus and Kollek, Ber. 1926, 59, [B], 1664). Diacetylene can also be prepared by the action of excess of alcoholic potassium hydroxide at 150-160° on butadiene tetrabromide (Lespieau and Prévost, Compt. rend 1925, 180, 675), and it has been obtained together with numerous other products by maintaining an electric are beneath the surface of ethyl alcohol (Müller, Helv. Chim. Acta, 1925, 8, 826).

Symmetrically disubstituted diacetylenes of the conjugated type R·C:C·C:CR can be smoothly prepared by the action of cupric chloride solution on the copper derivatives of monosubstituted acetylenes RC:CH (Straus and Kollek, l.c.), or by the action of iodine (2 atoms) on acetylenio Grignard reagents of the type CR: CMgX (Grignard and Tchéoufaki, Compt. rend. 1929, 188, 357). Symmetrical diacetylenic hydrocarbons of unconjugated type are obtained (1) by the action of acetylenyl bromides on magnesium (sometimes sodium) in ether,

Mg $2\mathsf{CR} \colon \mathsf{C} \cdot \mathsf{CH}_2 \mathsf{Br} \longrightarrow \mathsf{CR} \colon \mathsf{C} \cdot \mathsf{CH}_2 \cdot \mathsf{CH}_2 \cdot \mathsf{C} \colon \mathsf{CR},$

(2) by the action of monosodio-acetylenes on polymethylene bromides,

 $\begin{array}{c} 2R \cdot CCNa + Br \cdot [CH_2]_5 \cdot Br \\ \longrightarrow R \cdot C \cdot [CH_2]_5 \cdot C \cdot [CH_2]_5 \cdot C \cdot [CR,] \end{array}$

and (3) by the action of suitable dibromoolefins on polymethylene dibromides in presence of magnesium, followed by treatment of the product with alcoholio potassium hydroxide:

Br·[CH₂]₇·Br+2CH₂:CBr·CH₂Br

$$\begin{array}{c} \mathsf{Mg} \\ \longrightarrow \\ \mathsf{CH}_2 : \mathsf{CBr} \cdot [\mathsf{CH}_2]_{\mathfrak{g}} \cdot \mathsf{CBr} : \mathsf{CH}_2 \\ & \qquad \qquad \mathsf{KOH} \\ \longrightarrow \\ & \qquad \qquad \mathsf{CH} : \mathsf{C} \cdot [\mathsf{CH}_2]_{\mathfrak{g}} \cdot \mathsf{C} : \mathsf{CH} \end{array}$$

From those diacetylenio hydrocarbons which contain one or two replaceable hydrogen atoms in the molecule, alkyl derivatives can be obtained by the action of methyl sulphate or other suitable alkylating agent (see ACETYLENES) on a metallic derivative of the hydrocarbon:

$$\xrightarrow{\text{Me}_2\text{SO}_4} \quad \text{CMe}_{:}\text{C} \cdot (\text{CH}_2)_2 \cdot \text{C}_{:}\text{CH}, \text{ and } \\ \quad \text{CMe}_{:}\text{C} \cdot (\text{CH}_2)_2 \cdot \text{C}_{:}\text{CMe}$$

excess of hot alcoholic potassium hydrovide on the corresponding diolefin tetrabromides:

R CHBr-CHBr-[CH,]n-CHBr-CHBr-R'

R-C:C-[CH,]a-C:CR' Properties .- The diacetylenes are gases or houids, usually of pronounced odour. compounds which retain one or two reactive hydrogen atoms in the molecule react with ammoniacal cuprous chloride, alcoholie silver nitrate, etc., to give metallic derivatives in the same manner as does acetylene, and with capable of displaying additive properties similar hypohalous acid (or under certain conditions with halogens) to give halogeno-acetylenes Also all diacetylenes add chlorine, bromine,

iodine, or hydrogen halide at their unsaturated centres. Diacetylene, the simplest compound of the group, is a gas at room temperature (b.p. +95° to 10°/749 mm.; d0 0 7364, n0 1-43862). It readily yields explosive metallio derivatives and in liquid form polymerises easily to give darkcoloured solid products; when heated, it forms liquid polymendes. By the action of redine on the alver compound, or more conveniently, by the action of iodine and byposodous acid on the hydrocarbon, di iododiacetylene is obtained; impure specimens of the latter are violently explosive. Di-sododiacetylene is transformed by sodine in othereal solution to aabyoshexaiodobutadiene, m p. 1655-1665°. With iodine in chloroform solution discetylene yields the tetraiodide, C₄H₂I₄, m p. 58-59°; with bromine it reacts with unexpected slowness, the main product being a hexabromide, m.p. 183-5°, but it gives also some proportion of an isomeno bromide, m.p. 114°. Diacetylene capnot be regenerated satisfactorily from the hexabromide by the action of zinc m an alcohol or a cetone medium.

The conjugated diacetylenes can be bydro genated catalytically and can evidently display some tendency towards terminal addition since diphenyl diacetylene on dihydrogenation undergoes 1:4, 1:2.3 4-, and 1:1:4.4-addition simultaneously:

H. (CHPh.C:C:CHPh CPh:C·C:CPh CHPh:CH CH:CHPh CH,Ph-C:CCH,Ph

In general, each unsaturated centre of either conjugated or unconjugated diacetylenes is to those which characterise the simple acetylenes.

Conjugated Diacetylenes,

R C 'C C CH and R C C C CR. Methyl diacetylene (\(\triangle^{ay}\). Pentadi inene) 42° (c.) (A "Y-Hexadi-inene) 80" (c.) Ethyl n-Propyl-(△ay. Heptadi inene) (Δβδ.Hexadi-inene) Dimetbyl-130° (64°) {∆ye.Octadi inene) Diethyl-(A b Decadi Di-n-propyl-. 68°/12 mm. mene) Di-n-bntvl-(A en Dodecadi. 103°/8 mm mene) (A Ze Tetradecadi-Di-n amyl . 118°-119°/4 mm. mene) (αδ·Diphenyl·Δαγ. Diphenylbutadi-mone) (86 5-87") (αδ-Diphenyl-Δβδ. Dibenzylhexadi inenel (101°) Dr. B. phenylethyl discetylene (as Diphenyl-△ Yf-octadi-inene) (118°)

UNCONJUGATED BLACETYLENES,							
	Formula	B.p. (m p).	ď.	₩ _D			
Δ ^{αε} ·Heptadi-inene .	CH:C-[CH _{2]3} ·C:CMe	26°-27°/3 mm.	0 81021	1-4521*1			
Δαζ. ,, ,,	CH;C [CH,],C;CH	111 5°-112-5° mm.	0 816417	1-45117			
20 Octati mene	*** * * * * * * * * * * * * * * *	82°)10 mm.	D-3528M	1.48593			
$\Delta^{a\theta}$. Nonadi inene .	CH:C-[CH],-C:CH	55°-55 5°/13 mm. (-21°)	0 815921	1-45211			
Δ ακ. Undecadi-inene .	сн:с·[сн _т] ² -с:сн	82 5°-83°/2 mm. (-17°)	0 8182*1	1-45311			
Δ ^{αμ} ·Tridecadi-inene .	CHIC (CH.); C:CH	(-3° to -2°)	0 8262*1	1 454**			
△ L. Pentadecadi incne	C.H., C.C CH. C.C C.H.,	131°-135°/4 mm.	_	-			
△ do Hexadecadi inene	CH;C-(CH,),-C;CH	152°-155°/12 mm.		-			
Δζι_ ,, ,,	C,H,, C;C·CH, C;C C,H,	169°-170°/15 mm.	-	l –			
Δζκ " " "	C,H,1 C:C [CH,] C:C C,H,1	157°-158°/10 mm.		-			
Δηκ. Heptadecadı-inene		150°-155°/6 mm.		-			
Δηλ.Octadecadi-inene	C.H., C:C.[CH.], C:C.C.H.,	167°-168°/7 mm.		-			
Δ ar Eicosadi inene .	CH:C-[CH,], C:CH	(54°)		-			
△ Tricosadi inene .	C,H,,-C;C CH, C;C C,H,,	(-12° to -10°)	_	-			
△ * Tetracosadı inene		(38 5°-39°)	-				
	· · · · · · · · · · · · · · · · · · ·		1	H. F.			

DIAMOND.

DIACRIDINES v. ACRIDINE.

5:5-Diallylbarbituric acid, Allo-DIAL. barbitone, m.p. 171-172°. Hypnot London), B.P.C. (v. Barbiturio Acid). Hypnotic (Ciba,

DIALLAGE. An important rock-forming mineral belonging to the monoclinic series of the pyroxene group. It has the same composition as diopside and hedenbergite,

Ca (Mg, Fe) (SiO₃),

but with the addition of alumina passes into augite. It differs from the other members of the pyroxene group in possessing a fine lamellar structure, due to secondary twinning and partial alteration of the material. On the surfaces of foliation the mineral displays a shining metallic sheen, and on this account it is sometimes used as an ornamental stone. The colour is grey, brown or green. Diallage is of common occurrence as a constituent of gabbro.

L. J. S.

DIALLYL, $\triangle^{a\epsilon}$ -hexadiene (v. BUTADIENES AND POLYOLEFINES (non-conjugated).

DIALURIC ACID, Tartronylurea, v. BAR-BITURIC ACID (Vol. I, p. 625b).

DIAMOND. Carbon crystallised in the cubic system. The contrast presented by the physical characters of the two forms of crystallised carbon-diamond and graphite-is very striking. Diamond is the hardest of minerals and, indeed, of all known substances, but at the same time it is very brittle; on the other hand, graphite is one of the softest of minerals, and it is flexible and inelastic. The one is colourless and transparent and a had conductor of electricity, whilst the other is black and opaque and a good conductor. There is also a considerable difference in density—diamond, 3.52; graphite, 2.25.

The name diamond, in French and German Diamant, and similar in most other European languages, is a corruption of the Latin Adamas, which was used for this mineral by Manilius (A.D. 16) and Pliny (A.D. 100). This came from 'Αδάμας, meaning invincible, a word applied by the Greeks to hard metals and stones. The similar name adamantine-spar was nsed by the chemist Joseph Black for corundum. Although diamond and corundum are numbered respectively 10 and 9 in the mineralogist's scale of hardness, and with no other mineral falling between the two, yet, as shown by abrasive tests, diamond is 140 times harder than corundum, and more than a thousand times harder than quartz (No. 7 on the scale). It is on this high degree of hardness that all the technical applications of diamond depend; and a faceted gem never shows any signs of wear. The hardness is greater on an octahedron face than on a cube face, and on each face it varies with the crystallographic direction. Lapidaries assert that stones from Borneo and New South Wales are harder than those from other localities.

Crystalline Form.—Crystals of diamond are generally found singly and with faces developed on all sides; thus indicating that they grew freely from individual centres in some surrounding medium. By far the commonest form is the regular octahedron, the faces of which are often

"etched figures," whilst the edges are nearly always rounded and often marked with a pronounced furrow. Other forms are a hexakisoctahedron (six-faced octahedron), and less often the rhombic-dodecahedron and the cube, while orystals of distinctly tetrahedral habit are rare. Nevertheless, it is probable that crystals of diamond are really hemihedral (tetrahedral-cubic); the apparent octahedron with grooved edges consisting of two tetrahedra interpenetrating in twinned position. Distinctly formed twinned crystals are not uncommon. These are contact-twins with a face of the octahedron as twin-plane, and the two individuals flattened parallel to this plane, giving a triangular plate with re-entrant angles at the three corners. These are known to the Dutch cutters as " naadsteenen" (suture-stones). In addition to the rounded edges, noted above, the faces themselves are often curved, sometimes to such an extent that the crystals are almost spherical in form. This rounding, as well as the etched figures, suggests that the crystals have been corroded subsequent to their growth. An important crystallographic character of diamond is the existence of perfect cleavages in four directions parallel to the faces of the octahedron. (On the crystallography of diamond, v. A. Fersmann and V. Goldschmidt, "Der Diamant,"

Heidelberg, 1911).
Physical Characters.—In its optical characters, diamond is remarkable for its very high refractive and dispersive powers, the indices being: for red light (B) 2.40735, yellow (D) 2.41734, violet (H) 2.46476, and the dispersive value (H-B) 0.05741. The critical angles of total reflection corresponding with these indices are 24° 33′, 24° 26′, and 23° 56′ respectively. The angles between the facets of the brilliant form of cutting adopted for the diamond are so arranged that a ray of light entering the gem by the front facet will meet the back facets at a greater angle than the critical angle, and will be internally totally reflected, passing out again from the front of the stone, and with a chance that rays of different colours will be separated. A cut stone, therefore, shows much brilliancy or "fire," together with flashes of prismatic colours. Corresponding with the high refractive index, diamond displays a high degree of lustre, which is of the quality known as "ada-mantine" lustre. Uncut stones, especially when not quite clear, and with rough rounded surfaces, present a peculiar and very characteristic lead-grey metallic appearance. When pure, diamond is without colour and perfectly transparent. The best stones are described as of the "first water" or "blue-whites." More often there is a tinge of colour (" off-coloured " stones), usually pale yellow, brownish, grey or greenish. Stones of a good canary-yellow, coffee-brown, or black colour are not uncommon, but those of rich shades of green, blue, or red are very rare. The nature of the colouring matter is uncertain, but it is probably inorganic (e.g. iron and titanium in the brown stones). Changes in colour may be effected by the action of heat or by exposure to radium emanations. Crystals of diamond, being cubic, should be optically delicately marked with minute triangular pits or isotropic, but when examined under the polaris574 DIAMOND.

enclosures in the stone. This is due to a state of the surface has often been observed, but not of strain, but according to R. J. Sutton (1928) always confirmed. C. Doelter (1911), with stories of exploding crystals are mythical, crystals packed in charcoal or thorium and Luminescence (fluorescence and phosphorescence) is readily produced in some, but not all, diamonds by the action of sunlight, ultra-violet, Rontgen, cathode and radium rays, or simply by rubhing (triboluminescence). The cause of the lumines cence was attributed by G. F. Kunz in 1895 to a bydrocarhon which be called "tiffanyite." Diamond becomes positively electrified by friction. It is very transparent to the Röntgen raya; whilst glass imitations are opaque. Being a good conductor of heat, it can be distinguished from a glass imitation by touching with the tip of the tongue, diamond feeling much colder than glass; also a film of moisture from the breath will disappear more quickly off the surface of diamond than off glass The specific heat is less than that of graphits, and it increases rapidly with the temperature Colonriess diamonds (apart from those that are distinctly coloured) show amongst themselves differences in the absorption bands in the infra red and the quent presence of minute enclosures in diamond ultra violet; and some of them develop an electric current when exposed to ultra violet light-photoelectric effect (R Robertson, J J Fox and A. E Martin, Phil Trans 1934, 232A,

Chemical Characters.-The combustibility of diamond was first proved experimentally by the Florentine Academicians in 1694 Lavoisier, in 1772, found that the presence of air was necessary, and that "fixed air" resulted; but it was Smithson Tennant, in 1797, who first proved that equal weights of diamond and carbon yielded equal weights of carbon dioxide. More convincing experiments in this direction have These are crystalline aggregates, as distinct from been made by Davy, Dumas and Stas, A. single well-developed crystals. Krause (1890) and H Mosssan (1893). Powdered diamond hurns readily in air at a red heat Moissan determined the temperature of ignition around a common centre, giving rise to rounded In caygen to be 600-700°, according to J. Joly externer forms. These are consetting quite it is 850° in art. It burns with a small pale appeared ["hob-tuer" or "halls."] They blue flame, and in caygen is able to support are usually rough on the exterior, it being only its own combustion. It is infinishe even in the here that the fibres, or individual crystals. electric arc; and it can be heated at high have been free to develop crystal faces-in the temperatures (1,300) without siteration in interior these have interfered with each others carbon dioxide, however, at 1,200° the crystals are correded, with formation of carbon monoxide. It is unattacked by acids, caustie alkalis, iodic anhydride, hydrogen potassium sulphate, or a mixture of acdium chlorate and nitrie acid, but is oxidised when heated to 180-230° m a mixture of potassium dichromate and sulphuric acid. In fused potassium mitrate or sodium carbonate at about 900° amall atched figures are bonste at about 1907 annua accesse aguers are direction, a chance section of nor vall preced-alowly produced; at 1,2007 the actions is more both marinum and minimum values. The rapid, carbon dioxide being liberated. It is rounded pieces of bort are translucent to cloudy acted upon by aniphur vapour at 900°; and the of the opposition of the region of the combines with iron, yielding steel. Crystals are greasy to metallic lustre, and, usually, a peculiar teched by mother olvina (imageneum mitacis), leaden appearance; apgr. 3 50 In the trade The fact that diamond gives with oxidising

ing miccoscope they are frequently seen to possess isomerides. Observatious at high temperatures anomalous double refraction, especially around are somewhat contradictory; a mere blackening and heated to 2,500°, obtained only a superficial blackening without any alteration in the physical characters of the material. Moissan (1893). however, records the conversion into graphite at the temperature of the electric arc (about 3,600°); and Parsons and Swinton (1907) found that a dramond placed in the focus of cathode rays in escuo awelled up into a coka lika mass when the temperature reached 1,890°. In this connection the occurrence of cubes of graphite (cliftonite) in meteoric irons is aignificant; these are supposed to be paramorphs after diamond.

The asb remaining when diamond is burnt amounts, for colourless crystals, to 0 02-0 05%. but in the less pure carbonado it may reach 48% It consists mainly of iron oxide and aslaca, with some lime, magnesia, and titanium, The scon and titanium may represent the colouring matter present in the stone; but the freis not to be overlooked. In addition to liquid enclosures (carbon dioxide and water), the following materials have been recorded. black earbonaceous matter, magnetite, ilmenite or hæmatste, quartz, rutile, pyrite, gold, gamet, olivine, chlorite (?), apophyllite, etc. (R. J. Sutton, Min Mag. 1921, 19, 208, L. J. Spencer, shid 1924, 20, 245

Varieties .- Differing in certain points from the general characters enumerated above, there are two varieties of diamond-bort and carbonado-which, for technical purposes, are more important than the purer crystals used for gems.

Bort (boart ne boort) consists of a number of individual crystals with a radial grouping various gasea (hydrogen, mitrogen, aulphur growth, Although each individual possesses the dioxide and trioxide, and nitrous oxide) In cleavage characteriatio of diamond, yet for the whole mass there is no continuous cleavage For this reason, bort is tougher than the single crystals, which readily apht along the cleavage directions, and is thus better adapted for boring, turning and cutting tools. This, no doubt, is the explanation of the common atatement that bort is harder than diamond; further, aince in crystals the degree of hardness varies with the direction, a chance section of bort will present agents only carbon dioxide, whilst graphito those obtained by bruting), and all crystals and yields graphitic oxide, has led to the suggestion stones that are useless for gem purposes. Such that these two forms of carbon are chemical material is of value as an abrasive.

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and opaque, and often more or less porous, with somewhat the appearance of coke. It forms irregularly shaped masses, and consists of a confused aggregate of minute crystal grains of diamond. It therefore presents no cleavage as a whole, and is consequently tougher and less frangible than single crystals. It is less pure than ordinary diamond, yielding more ash (up to 48%) when burnt. An analysis of the ash gave Fe₂O₃ 53·3%, SiO₂ 33·1%, CaO 13·2%, MgO trace. The sp.gr. is less, 3·15–3·34 (the value for crystals being between the limits 3·502–3·525). The largest known piece of carbonado was found in Bahia, Brazil, in 1895. It weighed 631.9 g., that is, slightly more than the famous "Cullinan" diamond (which weighed 621.2 g.). Both carbonado and bort show considerable variations in texture, and with increasing coarseness of grain they may pass insensibly into ordinary diamond.

Occurrence.—India, from very remote times until the middle of the eighteenth century, was the only source of diamond (with the unimportant exception of Borneo). Since the discovery of the Brazilian and South African deposits the output has gradually fallen off, the fluctuating annual output being returned as only 20 carats in 1916 and 2,480 carats in 1934. The secondary deposits have, however, only been worked by native methods, and the original deposits have not been discovered. Indian diamonds being of the best gem-quality, there is thus a possibility of further developments. They occur associated with pebbles of jasper and vein-quartz in thin beds of conglomerate in the ancient sedimentary rocks (sandstones and shales) of the Vindhyan system (of pre-Cambrian age), which rest directly on the crystalline rocks. With the weathering of these strata, the diamonds are accumulated in the alluvial deposits of the present rivers. The districts where mining has been done fall into three main groups: a southern group in the basins of the Pennar, Kistna and Godavari rivers in Hyderabad and Madras; an eastern group in the Mahandi and Brahmani valleys in Bihar and Orissa; and a northern group between the Ken and Son rivers near Panna in Bundelkhand. A doubtful occurrence of diamond in pegmatite has been described from Wajra Karur near Bellary; and, unfortunately, nothing is known as to whence the diamonds found in the ancient sedimentary rocks were derived. (On Indian diamonds, v. Manual of the Geol. of India, vol. 3, Economic Geology, 1881, by V. Ball; vol. 4, Mineralogy, 1887, by F. R. Mallet; J. C. Brown, "India's Mineral Wealth,"

In Brazil, diamonds were discovered about 1725 in the alluvial gold-washings at Tejuco (now Diamantina) in Minas Geraes; although they have been since found in several other districts, notably in Bahia, this has remained the principal diamond-mining centre. Here, as in India, the diamond-bearing alluvial deposits have been derived from conglomerates

Carbonado, or "black diamond" (known in the trade as "carbon" or "carbonate"), is a finely granular crystalline material, dull, black which these were derived is not known. The Brazilian deposits are, however, remarkable for the variety of minerals found as well-worn pebbles ("favas") in association with the diamond; and the nature of these minerals, or indicators, has led to the suggestion that they, together with the diamonds, originally came from quartz-veins. (On Brazilian diamonds, see E. Hussak, "Os Satellites do Diamante," Rio de Janeiro, 1917; H. Preston, J. Soc. Arts, 1909, 58, 101; L. J. Moraes and D. Guimarães, Ann. Acad. Brasil. Sci. 1930, 2, 153, and Econ. Geol. 1931, 26, 502; C. W. Correns, Z. pr. Geol. 1932, 40, 161, 177).

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In South Africa, diamonds were first discovered in 1867 near Hopetown on the Orange River. In 1868 the important workings (" river diggings ") on the Vaal River were commenced; and in 1870-1871 the diamonds were traced to deposits of a unique type, in the neighbourhood of which the town of Kimberley in Griqualand West, Cape Province, very quickly sprang up. Here are situated the famous mines De Beers, Kimberley, Dutoitspan, Bultfontein and Wesselton. Other mines of the same type are the Jagersfontein and Koffyfontein mines in Orange Free State, and the great Premier mine (discovered in 1902) near Pretoria in the Transvaal. The diamond-bearing rock of these mines fills volcanic pipes, 200-300 yards (in the case of the Premier mine half a mile) across, penetrating vertically the surrounding beds of basalt, shale diabase, and quartzite, and extending to un-known depths. The rock, known as kimberlite or locally as "blue ground," consists mainly of a hydrated magnesium silicate with much the same composition as serpentine. It contains boulders and broken fragments of various rocks (basalt, shale, sandstone, quartzite, granite, mica-schist, eclogite, etc.) and minerals (ilmenite, pyrope, enstatite, chrome-diposide, pyrite, etc.) and is of the nature of a volcanic agglomerate. Diamond is present in this rock on an average of one part in fourteen millions (0.00008%), and in the richest part of the Kimberley mine one part in two millions. The diamond was brought up from below with the igneous material, which must have been derived from a basic magma rich in olivine. Although the origin of diamond has been the subject of much discussion, there seems no reason to doubt but that it orystallised from this basic magma. The rare presence of diamonds embedded in boulders of eclogite (a garnet-pyroxene rock) found in the blue ground has led to the suggestion that this is the mother-rock; but it is quite conceivable that the mineral crystallised from more than one kind of basic magma.

The early open workings ("dry diggings") at the Kimberley mines were soon replaced by a regular system of underground mining. hard blue ground, when brought to the surface, was formerly spread out on extensive floors and exposed to the action of the weather for about a year, when it is crumbly enough to enable the heavy minerals to be separated by washing. The present method is to pass the rock directly and sandstones (including the flexible sandstone through crushers and rollers. From the concenDIAMGND.

trato the diamonds are separated by means of a from some unknown source in Canada, at mechanical acreting table coated with grease, Murfreeabore on Arkansas asome damonds have to which the diamonds ashers, whist the other bone found in a serpentinued periodiste rock minerals pass over. Finally, the stones are discussed in a surpensitude of the cleaned by boting in a solution of caustic soda Econ. God. 1922, 17, 692).

The gradectors of diamond was at one time

South-West Africa, Southern Rhodesis, Belgian Congo and Tanganyka Territory; hat only a But now there are several competitors in other few of them have proved to be productave, regions, though South Africa still accounts for These pipes, of Cretaceous age, have no doubt more than half the world's total production, supplied the diamonds found in the sand-dunes In the following table is given the output for the in South-West Africa, and the rich deposits best and worst years since 1914. The total found in 1927 and 1929 along the coast respectionation in 1928 of 7,742,000 metric carats is tively south and north of the month of the equivalent to 1548-2 kg, or rather over 14 metric Orango River. Rich alinvial deposits were or long tons (5 metric carats=1 g), valued at discovered in 1926 in the Lichtenburg district £20,200,000. In 1936 and 1937 the production in south-western Transvani, where the gravels of reached eight and nine million earnts respecan ancient river system are richest in diamonds in tively, more than half in each year coming from large pot holes in the underlying dolomite rock. It is possible, however, that some of these alluvial diamonds may have been derived from rocks of much greater geological age, for small green diamonds are oceasionally found in the goldbearing conglomerate (" banket") at Modder-fontein in the eastern Witwatersrand. In these rocks, of pre-Cambrian age, the diamond must have come from some earlier source, of which no have come from some earlier source, of which no trace has yet been found. (On South Africas diamonds, see G. F. Williams, "The Diamond limes of South Africa," and ed, New York, 1907; F. A. Wagner, "The Diamond Fields of Southern Africa," Johannesburg, 1914; J. R. Sutton, "Damond—A Descriptive Treatise," London, 1923; A. F. Williams, "The Geness of the Diamond, "2 vols, London, 1932).

Since 1903 diamonds have been discovered in Since 1903 diamonds have been discovered in

several other parts of Africa. They have been found in conglomerates of Triassio age near the borders of Belgian Congo and Portuguese Angola and extending northwards into French Equatorial Africa (E. Polinard, Ann. Soc. Géol. Belg. Puhl. Congo Belge, 1929, 52, C179; 1934, 57, C65; Bull. Acad. Roy. Belg. 1931, 17, 137). In West Africa many small diamonds are found is of opecial interest. It was first observed in In West Africa many small dismonas are found is of epecial interest, it was first observed in allurial deposits in the Gold Coast, a few in the atone which field in September 4, 1888, near Liberra and Sierra Leone, and in 1934 a single the village of Nove-Ures, Nizhni Novgord, fine crystal of 10 carsts in Northern Nigera, Russis. This stone is composed of olivine In Tanganyida Tectairay a large kimberdes popt (875%) and large 1928 89%, logs/ther-with nuclear and the overlying gravel have been worked iron, trouble, chromitic, habe carbonaccous annea 1921 at Mabuki, 35 miles S.S.E. of Mwanza matter, and about 19% of damond, the last as well as the most habe of List Victorie, Savana and monate margine parish works. on the south shore of Lake Victoria. Several minuto grayish grains. As dull black grains it kimberlite pipes were found in 1926 farther has also been found in the meteoric stone of sonth near Shinyanga. Dismond is also Carcote, Chili. In several, though not in all,

with gold in alluvial deposits along the Mazarani in the medeons irons of Youndegin (Western and Pottor nivers. Finals of isolated crystal and Pottor nivers. Finals of isolated crystal and Australia, Smituhial (Fennessee), Smituh

Many kimberlite pipes are found over a wide practically a monopoly of South Africa, and by area in South Africa, some extending also into Belgian Congo.

DIAMOND PRODUCTION OF THE WORLD (in metric carats).

	1914	1922	1928	
Union of South Africa Leighin Congo Angola Cold Congo Cold Cold Cold Cold Cold Cold Cold Cold Cold	2,727,558 147,742 23,877 — 1,005 — 13,716 55 1,580	465,634 203,925 114,156 250,202 98,683 256 	2,254,649 2,114,675 603,142 1,647,700 237,511 696,826 64 24,681 41,865 214,474 607 824	
Total	2,916,000	1,337,000	7,742,000	

· Figures not available.

The presence of diamond in certain meteorites sonto near sampanga. Dismood is also Carcole, Claii. In several, though not in sil, reported from Bagamoyo on the east costs of the masses of meteoric rom from Cañoa opposite Zanzibar (E. O. Teale, Geol. Surrey: Dishlo in Arizona it has been found as colourles. The Armanyika Territory, 1931, Short Paper No. 1952, Start Sta

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in South Africa, in serpentinised peridotite in 1 Arkansas, and in diabase in New South Walcs; whilst in meteoric stones the matrix is also of the same general character. In these cases the diamond is very probably present as a primary mineral, and one of the first constituents to crystallise out from the basic silicate magma. In addition, we have the occurrence of diamond in the nickel-iron of meteorites. This may be regarded, as a magma of a still more basic type, and it is thus probable that diamond may crystallise from basic magmas of widely varying composition.

Many attempts have naturally been made to constrain the abundant element carbon to crystallise as the rare and valuable diamond. Considerable doubts have recently been exconsiderable doubts have recently been expressed as to whether diamond has ever been produced artificially (C. H. Desch, Nature, 1928, 121, 799; F. Krauss, "Synthetische Edelsteine," Berlin, 1929; F. Stöber, Chem. Erde, 1931, 6, 440; R. Brauns, Zentr. Min. A., 1931, 218; M. K. Hoffmann, Fortschr. Min. Krist. Petr. 1933, 18, 17). The alleged products have never been more than the minutest particles, and these have not been subjected to rigorous tests. The hardness test by scratching has been mainly relied on; but many carbides have a high degree of hardness, and spinel, which may crystallise as colcurless octahedra, readily scratches quartz. The appearance under the microscope alone is not reliable, and no definite determinations appear to have been made of the refractive index. specific gravity, or X-ray diffraction patterns.

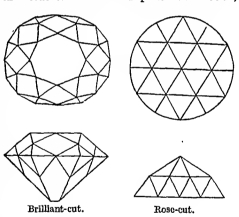
The experiments of I. Friedländer (1898), and of R. von Hasslinger (1902-1903), in which carbon was dissolved in molten magnesium silicate (olivine or kimberlite), bear some relation to the natural occurrences, but doubts have been expressed as to whether the microscopio crystals so obtained were really diamond. A method often tried is that first used by R. S. Marsden (Proc. Roy. Soc. Edin. 1881, 11, 20) and elaborated by Moissan (1893–1896; "Le Four électrique," Paris, 1897; and "The Electric Furnace," London, 1904), in which carbon is dissolved in silver or iron at a high temperature and the mass suddenly cooled by immersion in water or lead. A contracting crust is so formed and the molten interior subjected to great pressure. Sir C. A. Parsons (Bakerian Lecture, Phil. Trans. 1919, A., 220, 67; J. Inst. Metals, 1918, 20, 5), from a long series of experiments, however, concludes that pressure is net a necessary condition, and that the diamond had been formed from gases, probably carbon menoxide, occluded in the iron. E. de Boismenn ("Fabrication synthétique du Diamant," Paris, 1913) states that he obtained crystals up to 2 mm. across by the electrolysis of fused calcium carbide. O. Ruff (Z. anorg. Chem. 1917, 99, 73), in a systematic repetition of the various methods, obtained negative results, except perhaps by Moissan's method.

L. Duparc and P. Kovaleff (Compt. rend. Soc. Phys. Nat. Hist. Genève, 1924, 41, 108), from a consideration of the equilibrium relations of diamond and graphite, conclude that the stability of diamond increases with pressure and

was made to produce diamond at ordinary temperature under very high pressure, as in Spring's experiments. Carbon disulphide and a metal with a strong affinity for sulphur were subjected to a pressure of 8,000 atm. Dissolving the metal in acid, there remained a minute residue of colourless irregular grains which were birefringent with high refraction and which readily scratched glass. After some days the particles became covered with an opaque crust and fell to a fine powder. The suggestion that diamond was formed in this experiment is clearly not substantiated.

L. Sesta (Phil. Mag. 1929, 7, 488) repeated the experiments of M. La Rosa (1909-10). Using an intense intermittent electric are he obtained from sugar-charcoal spherical particles and minute tetrahedra with curved faces, with sp.gr. >3.2, H. >9, which burnt in oxygen, leaving no residue. M. K. Hoffmann (Zentr. Min. A, 1931, 214) found that a carbon electrode in liquid air with a current of 5,000 volts yielded glittering particles which resisted acids, but as the refractive index was less than 1.74 they were not diamond. J. Basset (J. Phys. Radium, 1934, 5, 471) by reducing carbides and various carbon compounds with metallic oxides under high pressure and temperature obtained only graphite.

Applications.—The value of diamond as a gom depends on its high degree of hardness enabling it to resist wear, and its high refractive and dispersive powers. These optical characters are, however, only brought into full play when the stone is faceted in a particular manner;



and for this reason, old stones cut by Indian lapidaries have often been re-cut. The form most suited for this purpose is the brilliant-cut. and stones so cut are known as brilliants. The rose-cut, or rosette, is only used for quite small stones usually mounted in number as a surround. The accompanying figures show a brilliant viewed on the large front facet, or table, and a side-view; and the rose-cut, viewed from above and the side. Being the hardest known material, diamond can only be worked very slowly and tediously with its own powder. The ground-form of the brilliant is approximately an octahedron, and this can be obtained, if necessary, diminishes with temperature, and an attempt by cleavage, or more safely by the modern

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method of sawing with a thin metal disc charged | absence of cleavage, are much more suitable

The large historical diamonds with their associations of mystery and romance are mostly of Indian origin; but these are now echipsed by the numerous much larger stones found in South Africa. The largest Indian atome still preserved is the "Koh-i noor," which was re-cut in England in 1852 as a brilliant of 106 th carats 1 (=21.786 g.). The largest Brazilian stone, the "Star of the South," found in 1853, weighed in the rough 254; "carata," and when cut 125 "carata." The largest known crystal of dismond is the "Cullnan," which was found in 1905 in the Fremier mine, Tranayasi This abowed a large eleavage surface, and was only a portion, probably about helf, of the complete crystal The portion as found weighed 621 2 g. (3,106 metric carats, or nearly I lb 6 ozs, a yourdupous). Being too large to cut as a single stone, it was cleaved and cut as nine large gems (weighing from 104 06 to 0 88 g) and ninety-six small brilliants, the yield of cut material amounting to 341%. The "Cullman," although the largest crystal, is not the largest piece of diamond that has hitherto been discovered A mass of carbonado weighing 631 9 g was found in 1895 in Bahia, Brazil.

As an ahrasive, diamond powder is the most effective agent available It is the only material with which diamond itself can be worked, and it is much used by lapidaries for cutting other hard stones. A thin disc of soft iron charged on the edge with diamond powder is used for sawing stones, both for ornamental purposes and in the preparation of thin microscope sections of rocks. The small steel tools used by engravers of cameos and intaglios, and also the ateel points used for boring precious stones, etc., are charged with dismond dust. The material used for abrasive purposes, though included in the trade under the term bort, consists of the waste material from shaping the better stones as brilliants, and also the less clear and spotted crystals not suitable for cutting as gems. It is crushed in a dismond-morter, splinters suitable for other purposes being picked out under a magnifying glass, and the remainder reduced to a fine powder.

The glazier's diamond is a crystal or fragment bounded by two rounded crystal-faces meeting in a curved or cusped cutting edge, For writing, drawing and engraving (e.g. fine scales and rulings on glass, metal, atones, ivory, etc.) fine splinters are used, or the diamond may be cut to a rounded point or chisel-edge. Diamond points are also used for drilling hard stones, glass, porcelain, teeth, etc. Carbonado and bort, by resson of their

¹ This was the old English carst of 203 409 mg A later value of the Anglish carst, as defined by the Board of Trade in 1893 and 1893, was 203 304 mg. The carstweight has now been standardised in all countries as the metric card of 200 mg (5 carsts-1 g). This has been the legal value of the carst in the British Isles. since April 1, 1914

with diamond dust. The brillaint is then than ordinary diamond for mounting as small roughly shaped out by rubbing two diamonds fragments (2-3 catats in weight) as cutting together -- a process known as bruting. The tools which may be subjected to shocks. These final grinding and polishing of the facets are are mounted in the crowns of rock drills, used performed on a rapidly revolving east-iron disc in tunnelling, mining, and sinking bore-holes for fed with diamond dust and olive-oil.

artesian wells, etc.; and on the edges of blade, band, circular, or wire saws for sawing large blocks of marble, granite, etc. Fragments, either in a rough form or fashioned to a suitable shape, are mounted as lathe tools for working stone, metal, etc. For example, for turning electric light carbons, the edges of watch glasses, finishing accurate turning work of the hard steel axles of instruments and machines of pre cision, boring cannon, forming the sharp edged furrows on the granding surfaces of millstones, for dressing emery wheels, etc.

Small cleavage plates or flat crystals of diamond drilled from each side with fine conical holes are used for drawing fine wire, down to 001 mm. in diameter. For the tantalum, tungsten and osmium filaments of electric lamps the metal is forced by pressure through the hole Bored diamonds are also used for the pivot supports of delicate instruments, such as chronometers, electric meters, etc

The epplication of diamond for optical purposes is limited by the high cost and difficulty of working, Diamond lenses would need to have a much slighter curvature than those of glass, and they would be immune from scratch-ing. Microscope objectives with such lanses were constructed by Pritchard under the direction of C R. Goring, in 1824-1826 A hemisphere of diamond would also increase consider-

ably the range and durability of refractometers References -In addition to the references quoted in the several sections above, the quofed in the several sections above, the following are given as heing of a more general character: M. Bauer, "Edulatemkunde," Lep-28, 1896—3 et de. by K. Schlossmacher, Leipzig, 1928-32—Eng. transl ("Frecous Stones"), by L. J. Spencer, London, 1904; C. Hintze, "Handbuch Bisso, vol 1.; C. Decker, "Lindbuch Bisso, vol 1.; C. Decker, "Handbuch Bisso, vol 1.; C. Decker, "Indone," Jondon, 1904; C. H. Sir W. Crookes, "Diamond," Jondon, "Jondon, "L. S. L. Sir W. Crookes, "Diamond," Jondon Bisso, vol 1.; L. A. Misra, "Diamond," stude, in Encycl. Brit. 11th & J. 1910; E. Boutan, "La Diamond," Plans, 1860; E. W. Streeter, "Tab Diament," Paris, 1866, E. W. Streeter, "The Great Diamonds of the World," London, 1882; W. R. Cattelle, "The Diamond," London, 1911; P. Grodzinski, "Diamant-Werkzeuge: Daratellung der Anwendung des Dismanten in Industrie und Gewerbe," Berlin, 1936.

o-DIANISIDINE, 4 4'-diamino-3 3'-dimethoxydiphenyl, colourless crystals, m.p. 135°, used as an intermediate for direct cotton dyes.

DIARSENOL. Canadian arsenical of the salvarsan type (J. Messner, Z. angew. Chem. 1917, 30, 49; Pharm. Monatab. 1921, 2,

DIASPIRIN, Trade name for the substance called succinyldisalicylic scid (OP. 196634) probably identical with the salicylyl succinide ether ester of H. P. Kaufmann (Z. angew. Chem. 1927, 40, 69); m p. 178°, antipyretic.

DIASPORE. A native hydrated alumina, Al,O, H,O or AlO(OH), crystallising in the orthorhombic system and isomorphous with goethite and manganite. It is usually found as scales or foliated masses with a perfect cleavage in one direction on which the lustre is pearly; but it is readily distinguished from other foliated minerals (mica, etc.) by its much greater hardness (H. 6½-7). Sp.gr. 3.4. When heated before the blowpipe it decrepitates violently, breaking up into pearly white scales (hence the name diaspore, from διασπείρειν, to scatter). In colour, it is usually grey or yellowish-brown, but pale-violet crystals are also found. It is usually found as an alteration product of corundum and emery, but occurs also as a mineral of contact-metamorphic origin in crystalline limestones. It is of importance as a constituent of bauxite, and in a particular type of bauxite, called diasporite, occurring in crystalline limestone in the Bihar Mountains, Hungary, it constitutes the bulk of the material.

L. J. S.

DIASPORITE v. DIASPOL

DIASTASE v. AMYLASE and BREWING.

DIATOMITE (Diatomaceous Earth, Infusorial Earth, Tripolite or Tripoli; Ger. Kieselguhr, Bergmehl). A form of opaline silica consisting of the siliceous remains of diatoms (Diatomaceæ). These are unicellular, free-swimming plants belonging to the large group of Algæ. The cell-wall, or frustule, of each minute individual consists of two shields or valves of siliceous material with elaborate and beautiful ornamentation. More than 10,000 species of diatoms have been described; they vary considerably in form—fusiform, elliptical, discoidal, globular and acicular. Material composed of individuals of the acicular form, the length of which varies from 0.093 to 0.156 mm. and the thickness 0.00156-0.00468 mm., is best suited for technical purposes. Diatoms multiply extremely rapidly and inhabit both fresh and salt water, being more abundant in cold latitudes. On death, their remains accumulate, under certain conditions, at the bottom of swamps, lakes or the open ocean. Besides being formed at the present day, such deposits are also met with in strata of Pleistocene and Tertiary age.

Diatomite is a loose, pulverulent material with a mealy or earthy texture, resembling soft chalk or dried clay in appearance. It is friable between the fingers with a harsh feel, and grates between the teeth. In colour it is white, grey, yellowish, brownish or greenish. It is very light, floating on water; sp.gr. of air-dried material 0.24 and of calcined 0.34. Its high degree of porosity depends on the fact that the shells of the diatoms are hollow. Good material absorbs four or five times its weight of water. Further, it is a bad conductor of heat, electricity and sound, and is resistant to heat and most chemicals. Of acids, only hydrofluoric attacks it, but it is slowly dissolved by a hot solution of caustic alkali. Chemically it consists of hydrated silica, mixed with various impurities, such as sand, hydroxides of iron, claycy, calcareous and organio matter. 70-90% of SiO₂ may be present.

Analysis I is of white material (dried at 105°) from Auxillac, Cantal, Auvergne; II, of green material from Auxillac; III, from Loch Cuithir (or Quire), Skye, Scotland; IV, from Richmond, Virginia; and V. of tripolite from Bilin, Bohemia (including also SO₃ 0·12, P₂O₅ 0·24, NH₄ 0·03).

	ı.	II.	III.	IV.	v.
	%	%	%	%	%
SiO ₂	91.61	88.56	88.73	75.86	74.20
TiO,	0.10	0.18	-	-	—
Al ₂ O ₃	1.53	$2 \cdot 16$		9∙88ๅ	6.81
Fe ₂ O ₃	2.22	0.73	0.67	2∙92∫	_
CaO	0.18	0.26	0.04	0.29	0.41
MgO	0.29	0.02	—)		-
K,0	trace	0.47	>	1.63	0.02
Na ₂ O	trace	0.44)		0.30
H,Õ	2.33	3.01	6.40	8.37	13.30
Organic matte	1.74	4.17	4.15	-	4.20

Tripolite (Ger. Tripel, Polierschiefer=polishing shale) differs from diatomite in being more compact with a laminated structure and consequently less light (sp.gr. 1.86) and porous. It merely represents the same material of a greater

geological (Tertiary) age.

The most extensively worked deposits of typical diatomite (Kieselguhr) are those on the Lüneburger Heide (heath) in Hanover. Here beds with a thickness of 10-15 m. occur in basins in Pleistocene sands. The uppermost layers are white, below they are grey, and at the bottom of the deposit green. The green material contains from 15 to 30% of organic matter, sufficient for calcination without addition of fuel. A bed of 5-7 m. covered by Tertiary clay and basalt is worked at Vogelsberg near Altenschlirf in Hesse. Similar deposits of recent to Miocene age are also widely distributed in the volcanic region of central France (Departments of Cantal and Puy-de-Dôme). Here the material is known as randannite and ceyssatite, from Randanne and Ceyssat respectively; and it is worked commercially under the name tellurine. (On the French and German deposits, see C. Schmidt, Ann. des Mines, Paris, 1910, 17, 370). Deposits are met with in poat-bogs at several places in Scotland, and beds up to 40 ft. in thickness of good quality material are worked at Loch Cuithir (or Quire) in the parish of Kilmuir in Skye (W. I. Macadam, Min. Mag. 1884, 6, 87; 1886, 7, 30, 35; 1889, 8, 135. Special Reports on the Mineral Resources of Great Britain, Mem. Geol. Survey, 1916, 5). Similar deposits also occur in Co. Antrim (J. H. Pollok, Sci. Proc. Roy. Dublin Soc. 1899, 9, 33). Extensive deposits, many of them of Tertiary age, are of wide distribution in the United States, especially in California (H. Ries, "Economic Geology," 6th ed., New York, 1930). Other localities are Monte Amiata in Tuscany, Victoria, Norway, Denmark, Hungary, Algeria, Chile, etc. The typical tripolite forms yellowish-white bands 0.5-1.5 m. thick in beds of clay, gypsum, and "Saugschiefer" (absorbent shale) in a Tertiary basin near Bilin in Bohemia.

In the preparation of the material for the

market, that of the purer white quality, after tates" C·N; OR; (3) a nitrogen atom in being dug out from open pits, is simply air- diszoamines (diszoamine compounds) dried in the open or in sheds. Organic matter is removed by calcination. Less pure material, containing sand, etc., is treated by washing and sedimentation. Iron is removed by the action of acids.

The varied uses of diatomita depend on its peculiar properties of porosity, hightness, nonconductivity, resistance to heat and acids. It was used by the Greeks and Romans for making light fire-proof tiles; and in the middle ages was mixed with meal for hread, under the impression that it had a medicinal value. As en absorbent it has been much used with nitroglycerine for the mannfacture of dynamite. The so-called dry sulphune acid is prepared by saturating calcined distomite with three or four times its weight of cid, Such material can be transported with little risk. Diatomito packed around carboys of acid will soak up leakage in case the vessels become cracked. It has also been used as an absorbent of homme, dis infectants and liquid manures. On account of its non-conductive properties it is used, sometimes with sebestos, as a light packing material for boilers, steam-pipes, cooking stoves, safes, refrigerators and refrigerating apparatus, fire proof and sound proof flooro and partitions of buildings, buildheads of ships Mixed with clay, lime, sawdust, etc., it can be used in the manufacture of fire-proof bricks and tiles; and having a low coefficient of expansion such material is specially adapted for making tubes, models, etc. It is used in the manufacture of various cements and artificial stones. Mixed with magnesia coment it gives a fire-proof coment for stoves; with shellac, ssaling-wax and gramophone records are made. It is also used as a filtering material for hquids and gases; and various liquids (wine, beer, acids, oil, syrups) can be clarified by stirring with 1% of distorate and allowing to settle. As a filler it is employed in the manufacture of rubber, cotton fabrics, paints, esfety matches, paper, papier-maché, soap, scouring-powders, tooth-powders, etc. Fused with calcium and magnesium borates it yields a non cracking glass for glazes and enamels, which can also be used for painting on glass. As a source of

Earth," Imp. Inst. London, 1928.)

OLAZINES v. AZINES. DIAZO- COMPOUNDS. These organic substances, like the azo- compounds, contain the of acdium natrite in 10-20% solution. characteristic group -Ng-, but whereas in the latter series this bivalent rodical is invariably attached to two carbon etoms, C.N.C. in the former it is generally, although not invariably, combined with only one carbon atom, the other valency being satisfied by: (1) an electro negative

C·N.·NHR

and diazoimines (azimino-compounds), which when derived from ortho-diamines have a cyclic configuration C.N. NH.C, whereas those from para-dismines are sometimes given a quinoneid structure N : CC:NH; (4) a sulphur atom in diazosulphonates C.N. SO,K, and diazosul-

phides C.N. S.N. C.
The diazocyanides C.N. CN contain the azogroup attached to two carbon atoms, one of which belongs to a cyanogen group, and these compounds which, in general, are distinctly coloured, form an intermediate group between the foregoing diazo- derivatives and the azocompounds.

Certain aliphatic diazo- compounds (v. snfra) contain the group -No- attached by two principal valencies to the same carbon atom.

L Diazonium Salts.

These diazo derivatives are generally, but not exclusively, prepared from aromatic amines, and the process, which is termed diazolisation, is carried out with the undissociated salt of the amine, and not with the free base. In a hydrolysing solvent like water, it is therefore custo mary to diazotise aniline and its denvatives with sodium nitrita (1 mol) and excess (21 to 3 equivalents) of mineral acid,

The eccentific and technical utility of the diazo reaction depends on the case and rapidity with which three hydrogen atome of the salt of an aromatic primary amine are replaced by nitrogen from nitrous acid to form the corresponding diazonium salt :

ArNH,CI+HO-NO=ArN,CI+2H,O

AROMATIC DIAZONIUM SALTS.

The diazetisation of the benzenoid and naphthalenoid amines is frequently carried out on a manufacturing scale owing to the employ. ment of aromatic diazonium salts in the production of azo- colonring matters. During the first decade of the present century more than edica, it is used in the proparation of codomi and such as the proparation of codomi and such as the proparation of codomi and as a such as the production of proparation of an abraive (g.r.) it, particularly the tupolic radiety in the production of proparation and anomaly in the production of proparation of an arrive, is used for publishing metals and stores of the partity replaced by a such as the sending the proparation of the production of proparation of partity replaced by a such as considerable when the production of the pr

The aromatic base is dissolved in dilute L. J. S. bydrochloric or sulphuric acid (3 equivalents in 10-20% aqueous solution) and the solution treated with an oquivalent quantity (1 mol) diazotisation is completed when the liquid rontams a slight excess of nitrous acid as indicated by starch and iodide paper.

With anthno and its homologues, the temperature of the diazo- solution is generally maintained at 0-5° to prevent, so far as possible, the decomposition of the unstable diazonium ion Ac in the diazonium salls C.N. Ac; (2) and the decomposition of the unstable diazonum oxygen atom in the diazohydroxides ("dazzo-hydroxides ("dazzo-hydroxides"), CN, OH and diazonoxides ("dazzo-hydroxides"), and the reases, e.g. the intranslation of the control of the con readily at 10°. The aminoanthraquinones are diazotised at 30-40° (Schaarschmidt, Ber. 1916,

49, 2678).

In alcoholic or acetic acid solution, diazotisation can be effected on the salt of the amine without using an excess of acid, and ethyl or amyl nitrite may, with advantage, be substituted for sodium nitrite. Gaseous methyl nitrite which has been employed in certain diazotisations (J.C.S. 1913, 103, 1398), is very readily prepared and is a cleaner reagent than amyl nitrite.

When treated with sodium nitrite, aniline hydrochloride in presence of excess of cold hydrochloric acid yields benzenediazonium chloride, whereas in absence of this excess of acid

it furnisbes diazoaminobenzene.

A product antecedent to either of these diazo-derivatives has been detected by Earl and Hall (J. Proc. Roy. Soc., New South Wales, 1933, 66, 455), who added I mol. of aniline hydrochloride in 40% aqueous solution at -8° to about 1.5 mols. of sodium nitrite in 50% aqueous solution covered with ether and cooled to -15°. A white precipitate separated and dissolved forthwith in the organic solvent. This ethereal solution was dried over anhydrous sodium sulphate and the ether removed under reduced pressure, when a mass of white needles separated consisting of aniline nitrite,

C₆H₅·NH₂,HNO₂

(yield about 60%). This salt becomes yellow at room temperature, and its aqueous solution

rapidly doposits diazoaminobenzene.

Benzenediazonium chloride, C₆H₅·N₂Cl, the simplest example of an aromatic diazonium salt, is prepared by adding the calculated quantity of amyl nitrite to aniline hydrochloride suspended in glacial acetic acid at 10°; it is precipitated from the clear solution thus obtained by the addition of ether.

Benzenediazonium sulphate,

C6H5·N2·HSO4,

is prepared in a similar manner from aniline sulpbate suspended in glacial acetic acid, the precipitation being effected with alcohol and

Benzenediazonium nitrate,

which was first precipitated from aqueous solution by ether and alcohol (Griess, Annalen, 1866. 137, 39), can also be produced in alcoholic or glacial acetic acid solution (Knoevenagel, Ber. 1890, 23, 2995; Bamberger, ibid. 1896, 29, 446; Hirsch, ibid. 1897, 30, 92; Hantzsch and Jochem, ibid. 1901, 34, 3337).

Diazonium salts have been obtained with a large variety of anions: platinichlorides, aurichlorides, stannichlorides, ferricyanides, nitro-prussiates (Ber. 1879, 12, 2119; 1885, 18, 965), chromates (Jahresberichte, 1867, 915; J.C.S. chromates (Jahresberichte, 1867, 915; J.C.S. phosphoric acid, for he obtained a small yield of 1905, 87, 1), carbonates and nitrites (ibid. 1905, 921), fluorides (Ber. 1903, 36, 2059), perchlorates (ibid. 1906, 39, 2713, 3146), picrates (ibid. 1906, 39, 2713, 3146), picrates (J.C.S. 1907, 91, 1316), azides (ibid. 1910, acetic acid solutions to nitrosyl sulphuric acid. 97, 1697; Ber. 1903, 36, 2056), thiosulphates,

bases, the diazotisation is carried out most | tungstates (J. Soc. Dyers and Col. 1901, 17, 279), thioacetates (Monatsh. 1907, 28, 247), borofluorides (G.P. 281055) (v. Vol. II, p. 43), phosphomolybdates and tungstates (J.C.S. 1909, 95, 1319), arylsulphinates (Ber. 1911, 44, 1415), zincichlorides, Rev. Gén. Mat. Col. 1914, 18, 4).

The zincichlorides are frequently used for stabilising diazo- compounds. Dianisidine hydrochloride is bisdiazotised in acid solution to which zinc chloride is added, when the crystalline double zinc salt (MeO·C₆H₃·N₂Cl)₂ZnCl₂ separates. A stabilised diazo- mixture is produced by mixing this zincichloride with partially dehydrated aluminium sulphate or sodium naphthalene tetrasulphonate and drying at 40° to 50° (B.P. 238676). 4-Aminodiphenylamine and its derivatives give stable diazonium zincichlorides and the technic variamine blue salt is of this type. technical

Diazonium chlorides added to a hydrochloric acid solution of lead tetrachloride furnish a yellow crystalline precipitate diazonium plumbichloride,

(ArN₂)₂[PbCl₆],

which on heating with alcohol gives the chloro-derivative ArCl (Chattaway, Garton and

Parkes, J.C.S. 1924, 125, 1981).

All diazonium salts are endothermic compounds and are liable to explode in the dry state (Berthelot and Vieille, Compt. rend. 1881, 92, 1076; Sventoslavsky, Ber. 1910, 43, 1497) and in some cases the explosion point has been determined (Ber. 1899, 32, 1691; 1900, 33, 527). The diazonium chromates and perchlorates decompose with considerable violence, and the former have been suggested as explosives (Bull. Soc. chim. 1867, ii, 7, 270; F.P. 73286).

Bisdiazonium salts (*Tetrazo- salts*). These compounds are produced from aromatic di-

amines.

When the two amino- groups are in the same aromatic nucleus, complete diazotisation is effected only with some difficulty. Metaphenylenediamine hydrochloride and sodium nitrite furnish the azo-colouring matters included under the term "Bismarck brown" (Ber. 1897, 30, 2111, 2899; 1900, 33, 2116), but when a solution of the base in excess of hydrochloric acid is added quickly to a large excess of well-cooled aqueous nitrous acid, complete diazotisation takes place and benzene-m-bisdiazonium chloride, C₆H₄[N₂·Cl]₂, is produced. When the nitrite is added quickly to the diamine hydrochloride, nitroso-m-phenylenediamine results. Para-phenylenediamine hydrochloride, when diazotised in the usual way, gives a mixture of diazonium and bisdiazonium chlorides, but when the hydrochloride is added to excess of nitrous acid, benzene-p-bisdiazonium chloride is obtained.

Schoutissen (J. Amer. Chem. Soc. 1935, 55, 4535) has obtained evidence of the diazotisability of o-phenylenediamine by treating the diamine in sulphuric acid with nitrosylsulphuric acid and phosphoric acid, for he obtained a small yield of o diiodobenzene on adding potassium iodide. Hodgson and Walker have bisdiazotised all o-dichlorobenzene (J.C.S 1935, 530).

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dismine is diszotised with difficulty, owing to its acid, sithough this coincidence is merely oxidisability, but in producing its azo- derivatives this difficulty is surmounted by working J.CS. 1923, 123, 228). with its monoacyl- derivatives and diagonisms. In an analytical at in two stages with an intervening hydrolysis (B.P. 18783, 1891). The heteronucleal naphthylenediamines, excepting the 1.8- compound, can be completely distortised, yielding naphthalene-bisdistronium salts (Ewer and Pick, G.P. 45549, 45788; Badische Anilin und Soda Fabrik, G P. 130475; Morgan and Micklethwait, J.CS. 1910, 97, 2558). Benzidine, tolidine, diameidine, and other similar heteronuclear diamines of the diphenyl and ditolyl series are readily diazotised, furnishing hisdiazonium salts (e.g. diphenyl-bisdiazonium chlorids, Cl N2 C4H4 C4H4 N2 C1), which are employed in the manufacture of substantivo azo- dyes.

The upper limit of diszotisability in the benzene series has been determined by a study of trammomesitylene (I), which when treated with nitrosyl sulphate in atrong sulphurio acid gives indezole derivatives. However, diazotisation of this trismino was successfully effected in aqueous hydrochlorio acid, when even with a large excess of soid and nitrite only two of the three amino groups became diazotised concurrently, and this twofold diszotisation was demonstrated by isolation of ammomesitylenebisdiszonium aurichloride (II) and aminomesitylene bisazoimide (bis-triazomesidine III). The latter compound was produced by adding sodium szide to the diazo. solution.

The non-diazotisability of the third amino-

tained from o phenylenediamine a 70% yield of gives tristriazomesitylene (IV), a compound having the remarkable molecular formula In the naphthalene series, 1 4-naphthylene- HaCoNe, which suggests a polymeride of prussic numerical and not chemical (Morgan and Davies,

In an analytical atudy of the titration of aromatic amines by the diazo- reaction it was found that 1.3 5-traminobenzene in hydrochloric acid solution gave figures corresponding with threefold diazotisation; 2,46-triaminochlorobenzene was completely diazotised only in alcohoho solution (Phillips and Lowy, Ind. Eng. Chem. [Anal.], 1937, 29, 381). VELOCITY OF DIAZOTISATION.—Under the

usual experimental conditions, diazotisation proceeds with great rapidity, but in N/1000 solutions, the velocity of this reaction has been determined by a colorimetric incthed for estimat. ing the free nitrous acid. The reaction is of the second order, and the values of the constant

 $\frac{1}{t(a-x)}$ are 0.036 for aniline, 0.038 for p toluidine, and 0 041 for m-xylidine, with one molecular proportion of free acid at 0°. At higher temperatures the rate is increased (Hantzach and Schimann, Ber 1893, 32, 1991; 1900, 23, 527; Tassully, Compt rend, 1913, 157, 1148; 1914, 158, 335, 489). Both colorimetric and electrolytic determinations of the velocity of diazotisetion show that, in the absence of disturbing influences, all the aromatic amines are diazetised at the same rate

ALTERNATIVE METHODS OF PRODUCING ARO-MATIO DIAZONIUM SALTS - Netrosyl chloride or MATIO DIADOPTO ALTYS — NIVER CHOPME WE be borned in the born suggested as a discrete promote has been suggested as a discrete promote the beautiful promote the suggested of the suggested and Kenson Amer. Chem. J. 1835 59; Kaatle and Kenson Amer. Chem. J. 1840 17, 91). Mirroyl sulphate has been used the advantage (Patet and Girard, D.R.P. 6034; Ber 1870, 12, 365; Morgan and Macklethus, J.C.S. 1910, 97, 2559). This reagent or sodium mirrted desiphers and discrete displayment and the support of the employed in diazotising alightly basic amines (Morgan and Evens, J.C.S. 1919, 116, 1130; cf. J. pr. Chem. 1906, [u], 74, 275). Barum situte, when employed in conjunction with aulphuric acid, furnishes diazo- aclutions free from inorganic salts (Witt, Ber. 1903, 38, 4388). The last-named investigator has brought forward an interesting process which consists in dissolving the base in cold concentrated miric acid and adding the calculated amount of potassiam metabisulphite required to produce aufficient nitrous acid for the diazotisation (Ber. 1909, 42, 2953).

For unsubstituted aromatic amines the direct method of diazotisation by adding nitrite to the cold acid solutions is generally efficacious, but for amines containing nitro, carboxyl, or sulphome groups the inverted method of mixing the amine with nitrite and running this mixture in cold acid is preferable. Highly aubstituted amines, such as 4 chloro 26dinitraniline and pieramide (2 4 6-trinitroaniline) group is not due to sterie hindrance, for when are dissolved in plansi action add and treated the bisduncarium radicals have been replaced with untroyslupiduric acid (Missim, Herty two transe groups the resulting histriano-Chim Acta, 1920, 3, 626; Blancy, tod., mesdine can still be diarotised, and on further 1925, 8, 780). Schoutissen dissolved highly treatment with sodium arise its diazonium sail best substituted bases in authoria acid and added a acid. Diazotisation was promoted by diluting the solution with phosphoric acid (sp.gr. 1.70) at 0° (J. Amer. Chem. Soc. 1933, 55, 4531; Rec. trav. chim. 1935, 54, 97). Similar methods are used to diazotise amines of the anthraquinone series (J. pr. Chem. 1906, [ii], 74, 275; Ber. 1904,

37, 4185).

In the diazotisation of 2:4-dinitro-a-naphthylamine the process is usually attended by the climination of the ortho-nitro- group, and a similar change occurs in the case of certain dinitroanilines. In order to avoid this formation of diazo- oxides it is suggested to diazotise such feehlo bases in cold acetic acid with a solution of sodium nitrite in concentrated sulphuric acid. Even 2:4:6-trinitroaniline (picramide) can he

diazotised by this process.

This mode of diazotisation may also he employed in converting 2:4:6-trinitro-m-pheny-lenediamine into its hisdiazonium salt (Misslin, Helv. Chim. Acta, 1920, 3, 626; Meyer and Tochtermann, Ber. 1921, 54, [B], 2283).

Diazonium hromides are formed by the interaction of hydrazines and diazonium perbromides

$$2R \cdot N_2Br_3 + RN_2H_3 = 3R \cdot N_2 \cdot Br + 3HBr$$

(Chattaway, J.C.S. 1908, 93, 958). An electrolytic process for the production of diazonium salts has been patented (Boehringer & Sons, D.R.P. 152926, and B.P. 2608 of 1904).

Diazonium salts can also be prepared from compounds not containing amino- groups; mercury diphenyl and nitrous fumes give benzenediazonium nitrate (Ber. 1897, 30, 509), and the same salt is produced from nitrosobenzene and nitric oxido in chloroform solution (ibid. 1898, 31, 1528).

Aniline nitrate is transformed into benzene diazonium chloride by zinc and hydrochloric acid (Möhlau, Ber. 1883, 16, 3080) or by titanous chloride (Knecht, J.C.S. 1924, 125, 1537).

Certain complex or highly substituted amines resist diazotisation unless treated under pressure (D.R.P. 143450) or with excess of nitrous and mineral acids (Annalen, 1873, 165, 187; 1891, 260, 224; Ber. 1900, 33, 520).

(For indirect methods of forming diazonium salts, v. Annalen, 1879, 199, 320; 1888, 243, 282; 1892, 270, 116; Ber. 1893, 26, 2190; 1899, 32, 1809; 1900, 33, 1718; J. pr. Chem. 1896, [ii], 54, 496; Gazzetta, 1900, 30, 1526; Bull. Soc. chim. 1891, [iii], 6, 94.)

Fuming nitric acid reverses the coupling process and regenerates the diazonium salt from its azo- derivatives (Meldola, J.C.S. 1889, 55, 608; 1894, 65, 841; O. Schmidt, Ber. 1905, 38, 3201). Diazonium chlorides are regenerated from azo-dyes by decomposing the latter with chlorine or hypochlorous acid (J. pr. Chem. 1912, [ii], **85**, 325.

REACTIONS OF AROMATIC DIAZONIUM SALTS.

In addition to their power of coupling with phenols, aromatic amines and their sulphonic acids to form the technically important azocolouring matters, the diazonium salts are of the hydroxy-derivative with a portion of the synthetical agents of the greatest value, and the still undecomposed diazonium salt (J.C.S.

solution of nitrosyl sulphuric acid in the same | following is a synopsis of the principal reactions in which they take part:

> 1. Decomposition of Diazonium Salts by Water.

> (v. Ber. 1874, 7, 1061; 1894, 17, 2704; 1899, 32, 1773; 1890, 23, 3635; 1902, 35, 89; J. pr. Chem. 1881, [ii], 24, 476; J.C.S. 1902, 81, 77; 1903, 83, 688; 1905, 87, 5; 1906, 89, 19).

The decomposition of soluble diazonium salts hy water is a unimolecular reaction, and is represented by the well-known equation:

$$k = \frac{1}{t} \log_e \frac{a}{a - x}$$

The relative stability of the diazonium salts is ascertained by comparing the values of the constant k for a given temperature. The most successful method of making this determination consists in heating solutions of the diazo- compounds at a fixed temperature, and measuring the rate at which nitrogen is evolved.

These quantitative decompositions show that the diazonium salts from aniline and orthoand meta-toluidine are among the least stable of the series, whereas considerably greater stability is exhibited by the diazotised nitrani-lines and the diazonium sulphonates derived from the benzenoid and naphthalenoid aminesulphonic acids.

The rate of decomposition of diazo- compounds increases with tho temperature and in accordance with Arrhenius's formula for the temperature coefficient (v. CHEMICAL AFFINITY,

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$$\log_e \frac{k_1}{k_2} = \frac{E}{R} \left(\frac{1}{T_2} - \frac{1}{T_1} \right)$$

The rate of decomposition is independent of the quantity of mineral acid present, and equivalent solutions of benzenediazonium chloride, bromide, sulphate, nitrate, and oxalate decompose with the same velocity; this change is, however, accelerated by colloidal platinum or silver, owing to catalytic action (Hantzsch, Ber. 1900, 33, 2517; Hausser and Muller, Bull. Soc. 1900, 33, 2517; Hausser and Muller, Bull. Soc. chim. 1892, [iii], 7, 721; 1893, 9, 353; Compt. rend. 1892, 114, 549, 669, 760, 1438; Schwalhe, Z. Farb. Ind. 1905, 4, 433; Ber. 1905, 38, 2196, 3071; 1909, 42, 196; Cain, J.C.S. 1902, 81, 1412; 1903, 83, 206, 470; 1905, 38, 2511; Euler, Annalen, 1902, 325, 292; Veley and Cain, Trans Faraday Soc. 1909, 5, 1) Trans. Faraday Soc. 1909, 5, 1).

In general, the decomposition of a diazonium salt by water leads to the formation of a phenol:

$$R \cdot N_2 \cdot HSO_4' + H_2O = R \cdot OH + H_2SO_4 + N_2$$

The sulphate is the hest salt to employ in this reaction, because the chloride and nitrate are apt to lead respectively to alternative or secondary changes. Benzenediazonium sulphate and its homologues furnish phenol and the cresols and xylenols; diphenyl-4:4'-hisdiazonium sulphate gives rise to 4:4'-dihydroxydiphenyl. Aurin is similarly obtained from diazotised rosaniline (Annalen, 1878, 194, 306).

In certain cases, more especially in the naphthalene series, the product is contaminated by azo- compounds, resulting from the coupling 1903, 83, 221). To ohviate this difficulty, so far as possible, excess of scid is used in the decomposition. The manufacture of the gnaphtholsulphonic acids (e g. OH: SO-H=1:4 and 1:8) from the corresponding a-naphthyl-ammesulphonic acids furnishes typical examples of the technical application of the process. When the phenolic product is volstile in steam, the normal change is facilitated by operating with solutions boiling above 100°. A good yield of guaiacol is obtained by adding the diago- salt of ortho anisidine to a mixture of sodium sulphate and ddute sulphuric acid boiling at 135-145°, the volatila product being removed at the moment of formation (Kalle & Co , B.P. 7233 nf 1897; cf. Annalen, 1889, 253, 281; Ber. 1874, 7, 1008; D.R.P. 167211; J.C.S. 1906, 89, 19; cf. Willstätter and Schndel, Ber. 1918, 51, 782).

Secondary reactions occur even in the aim-plest cases where the boiling is protracted (Ber.

dazo compounds by water, v. Ber 1874, 7, 1061, groups by the reducing action of ethyl alcohol 1884, 17, 2704; 1899, 32, 1773; 1902, 35, 89, has been applied successfully to a series of J.C.S. 1898, 89, 1327; 1902, 81, 77; 1905, 87, 5,

1906, 89, 19).

The disze compounds are very sensitive to light, which induces the phenolic decomposition even in certain cases where the action of hot water or ands is very alight (J.C.S. 1907, 91, 35; cf. Compt. rand. 1881, 92, 1074; Ber. 1890, 23, 3131; 1901, 34, 1668; D R.P. 53455, 56606, J.S.C.I. 1890, 9, 1001).

2. Decomposition of Diazonium Salts by Alcohols.

The reaction between diazonium salts and alcohols may proceed in two different ways:

I. R.N.X+C.H.OH

The former of these changes was originally regarded as the general reaction, but more recent researches have shown that the second is the normal change. The factors determining the course of the reaction are, however, numerous and complicated, involving (1) the nature of the alcohol; (2) the influence of substituents in the diazonium salt; (3) the preseure and temperature at which the decomposition is affected.

The tendency to replace the diszo- group by hydrogen increases with the molecular weight of the alcohol employed. Benzenediazonium chloride or sulphate and methyl slcobol give amsole exclusively. With ethyl alcohol, these salts yield chiefly phenetole, but benzen is also pro-duced. The diazonium sulphates from p chloro-and p brome entina furnish nnly p-chloro- and p brome benzene with ethyl alcohol, but with methyl alcohol they give rise mainly to the methyl ethers of p-chloro- and p hromo-pheuol (Amer. Chem. J. 1898, 20, 229; cf. Annalen, 1871, 159, 191; Ber. 1901, 34, 3337; 1903, 36, 2061: Amer. Chem. J. 1893, 15, 518: 1897, 19, 531, 547, 561).

The drazo- salts of the simpler amines (anshipe and its homologues) react chiefly in the direction indicated by the second of the fore. going equations, and yield mixed others, but the presence of acid aubstituents

(CI, Br, NO, CO,H, NH SO,R)

favours the predominance of the first change, The nitranilines (ortho-, meta-, and para-) give respectively 87, 51, and 40% of nitrobenzene, the 1:2-, 2:1-, and 1-4-nitronaphthylamines furnish the corresponding nitronaphthalenes, whereas a- and \$ naphthylamines yield a- and B ethoxynaphthalenes (Remsen and others, Ber. 1885, 13, 65; Amer. Chem. J. 1880, 8, 243; 1887, 9, 587; 1891, 13, 153; 1892, 14, 15).

The yield of alkyl ether increases with rise nf pressure, but the addition of zinc dnst or an alkalı (sodının hydroxide or ethoxide, potassium

heterocyclic diamines, tha diamino-cyclic 1:3diazalines, of which the following base (formula I) is the simplest example. This diamina and its analogues are diazotised by Schoutissen's method with nitrosyl sulphurio acid in concentrated phosphone seid and after dilution the soluble bisdiazonium salt is reduced in solution by boiling alcohol. The resulting cyclic I 3-diazalme (formula II) is extracted with other (Morgan and Stewart, J.C.S. 1938, 1292).

NH.

IJ. 3. Replacement of the Olazo- Group by

Hydrogen. In addition to the action of alcohol, the diazo group can be eliminated by the use of a

variety of reducing agents:

(1) The diazonium salt is reduced to the correspouding hydrazine and the hydrazine group NH-NH, removed by boiling with copper sulphate (Baeyer and Pfitzinger, Ber. 1885, 18, 90,

(u) Reduction of the diazo compound with alkaline sodium stannite (Friedländer, Ber. 1989. 22, 587)

(iii) By various other reducing agents, stannous formate (Ber. 1885, 18, 357; J. pr. Chem. 1889, [in], 40, 97), hypophosphorous acid (Ber. 1902, 35, 162), sodium hyposulphile (ibid.

1907, 40, 858), formic acid and copper powder [(ibid. 1890, 23, 1632), aluminium powder in alcohol (Morgan and Evens, l.c.).

4. Replacement of the Diazo- Group by Halogen, Cyanide, Cyanate, and Thio-

cyanate Radicals.

Fluorine may he introduced into the aromatic nucleus in place of the diazo- group hy treating the diazonium salt with hydrofluoric acid. Diazoamino- compounds (v. infra) are also employed in bringing about this replacement, and when treated with concentrated hydrofluoric acid they yield the corresponding fluoroderivative; fluorohenzene is thus produced diazoaminobenzene or diazobenzenefrom piperidide (J. pr. Chem. 1870, [ii], 1, 395; Ber. 1875, 8, 893; 1879, 12, 581; 1889, 22, 1846).

A general and more convenient method of

introducing fluorine into the aromatic nucleus consists in adding 40% horofluoric acid to the diazo-solution from the aromatic hase when the corresponding aryldiazonium horofluoride is precipitated. These borofluorides are generally very stable salts; they are not explosive and may be crystallised from warm water. When a diazonium borofluoride is gently heated it evolves nitrogen and boron trifluoride, whilst fluorine replaces nitrogen in the aromatic ring. Thus benzenediazonium horofluoride (decomposing at 121-122°) furnishes fluorobenzene, b.p. 85°/760 mm. Toluene-p-diazonium borofluoride (decomp. 110°), m-xylene-4-diazonium borofluoride (decomp. 108°), and naphthalene-adiazonium borofluoride (decomp. 113°) give respectively p-fluorotoluene, 4-fluoro-m-xylene, b.p. 143-144°, and a-fluoronaphthalene, b.p. 110°/17 mm.

The bisdiazonium salts from benzidine yield the corresponding bisdiazonium borofluoride (95% yield, decomp. 137-138°), and this salt is pyrolysed into 4:4'-difluorodiphenyl (b.p. 115-

116°/11 mm., 80% yield).

m- and p-Phenylenediamines are similarly converted into bisdiazonium borofluorides which yield respectively m-diffnorobenzene, h.p. 82-83°/752 mm., m.p. -59°, and p-diffuorobenzene, b.p. 88·5°/767 mm., m.p. -13° (Balz and Schiemann, Ber. 1927, 60, [B], 1186; Schiemann and Bolstad, ibid. 1928, 61, [B], 1403; cf. ibid. 1929, 62, [B], 1794, 1805; 1930, 63, [B], 3035). p-Nitrobenzenediazonium borofluoride decomposes smoothly into p-fluoronitrobenzene.

The first really successful process for replacing the diazo- group hy chlorine (or bromine) was discovered by Sandmeyer; it consists in adding the diazotised solution to cuprous chloride (or bromide) dissolved in warm hydrochloric (or hydrobromic) acid, when nitrogen is evolved and the corresponding chloro- or hromoderivative is produced (Ber. 1884, 17, 1633, 2050; 1885, **18**, 1492, 1496; 1886, **19**, 810; 1890, 23, 1628, 1880; Annalen, 1893, 273, 141)

The first stage in the Sandmeyer reaction is considered to he the formation of an additive compound of diazonium salt and cuprous chloride; the decomposition of this inter-mediate product proceeds readily only above a

141; cf. Votoček, Chem. Zeit. Rep. 1896, 20, 70; Hantzsch and Blagden, Ber. 1900, 33, 2544). The substitution of bromine for diazonitrogen follows a similar course (Phil. Trans, 1864, 154, 673; Annalen, 1866, 137, 49; Amer. Chem. J. 1891, 13, 486; Gazzetta, 1890, 20,

The foregoing substitutions are also effected hy the addition to solutions of diazonium chlorides or bromides of finely divided copper or the copper bronze of commerce (Gattermann, Ber. 1890, 23, 1218; 1892, 25, 1074; Ullmann, ibid. 1896, 29, 1878; cf. Angeli, ibid. 1891, 24,

Iodine is readily substituted for the diazocomplex by treating the acid diazo- solutions with potassium iodide even in the absence of

copper or cuprous salts.

As an alternative to the Sandmeyer and Gattermann reactions, the replacement of a diazonium group by chlorine or hromine is effected hy converting the diazonium salt either into mercurichloride hy adding mercuric chloride in aqueous potassium chloride or into mercurihromide hy adding mercuric hromide or nitrate in aqueous potassium hromide. The double diazonium mercurihalide is dried by washing with acetone, methyl alcohol or ether, mixed with twice its weight of potassium chloride or bromide and heated until the diazo- compound is completely decomposed. This procedure is effective in the following cases: 2:2'-diaminodiphenyl (2:2'-xenylenediamine) is converted into 2:2'-dichlorodiphenyl or 2:2'-dibromodiphenyl (yield 80%); p-bromobenzenediazonium bromide with mercuric bromide and potassium bromide gives p-dibromobenzene, and β -naphthylamine is converted similarly into 2-bromonaphthalene (yield 65%). The double diazonium salts with antimony trichloride and stannic chloride may be utilised in a similar manner (Schwechten, Ber. 1932, 65, [B], 1605).

Benzenediazonium chloride when treated with potassium and mercuric iodides in methyl

alcoholic solutions yields successively

C₆H₅·N₂·Hgl₃ and (C₆H₅N₂)₂Hgl₄. are vellow compounds decomposing h are yellow compounds de 63–64° and 65° respectively. Both \mathbf{W} hen maintained at the ordinary temperature these mercuric iodides pass mainly into iodobenzene and mercuric iodide with loss of nitrogen, hut in addition the former gives a 10% and the latter a 3% yield of diphenyliodonium tri-iodomercuriate $[(\tilde{C}_6H_5)_2[]H\tilde{g}I_3$, yellow crystals, m.p. 171.8° (Nesmejanov, Z. anorg. Chem. 1929, 178, 300).

The Sandmeyer reaction includes the important synthetical operation of replacing the diazo- group by a cyanide radical, the diazonium salt being added to a warm solution of potassium cuprocyanide. The Gattermann process is also applicable to this synthesis, the addition of copper powder to a mixture of diazonium salt and potassium cyanide inducing a similar replacement. The cyanate radical is introduced in the place of the diazo- group hy adding successively to a solution of diazonium salt, potassinm cyanate and copper powder (Gattermann, Ber. 1890, 23, 1218; certain temperature, which is different for each 25, 1086). The thiocyanale group is introduced diazonium salt (Erdmann, Annalen, 1893, 272, hy adding potassium thiocyanate and cuprous

throcyenate to a solution of diazonium sulphate | and Storch, Ber. 1893, 28, 471). The addition (Cattermann and Haussknecht, Ber. 1890, 23, of hydroxylamine to a solution of diazonium

738 ; Thurnauer, toid. 770).

Although cuprous halidea are employed in the Sandmayer reaction for replacement of diazo group by chlorine or bromme, nickel cyanide is better than cuprous cyanide for the introduction of cyanogen, and ferrie salts are the best catalysts for introduction of the thiocyanogen (CNS) group (Compt. rend. 1926, 183, 421).

5. Replacement of the Diazo. Group by Sulphur and Sulphur-containing Radicals, These reactions are of importanto as furnishing methods for the synthetical production of

sulphonic acids.

Thiophenols or disulphides are obtained by treating diazonium salts with potassium xanthate and then hydrolyang the resulting aromatic xanthate (Leuckart, J. pr. Chem. 1890, [11], 41, 179; cf. Klason, Ber. 1887, 20, 349; Walter, Chem. Soc, Proc. 1895, 11, 141). Phenyl sulphide is produced by treating benzenediazonium ealts with hydrogen sulphide or ammonium eulphide (Graebe and Mann, Ber 1882, 15, 1683). Benzenoid diazonium salts yield sulphides on treatment with aodium cuprous thiosulphate, but diszotised a naphthylamme only a-azonaphthalene (Bornstein, Ber. 1901, 34,

The replacement of a diazonium by a sul phinic group is effected by saturating with sulphur dioxide an acid solution of diazonium sulphate and adding copper powder; nitrogen is evolved and the resulting sulphinic acid ex tracted with ether (Gattermann, Ber 1899, 32, 651) 1136; BP. 26139 of 1898; DRP 95830)

The foregoing throphenois, disulphides and sulphinic acids yield the corresponding sulphonic acids on oxidation (D.R.P 70286; BP 11865 of

1892). 6 Replacement of the Diazo- Group by

Nitro-, Nitroso-, and Amino- Radicals. The introduction of a nitro- group is brought about either by heating the diazonium salt with nitrous acki and cuprous oxide or by mixing the diazonium sulphate with cupro-cupric sulphite and adding excess of alkali nitrite latter procass, 246-tribromobenzenediszonium sulphate gives 65% of 1 nitro 2 48 tribromo benzene and diazotised \$ naphthylamme gives and Wormall, J.CS 1921, 119, 2088) 25% of β nitronaphthalene, a substance prepared with considerabla difficulty by other 460; Amer. Chem J. 1888, 10, 368, Ber. 1902, processes (Sandmeyer, Ber. 1887, 20, 1917; 25, 1964. The Tey the replacement of the diazo group by diazonium borofluorides have been utilised in arsune and atibinic complexes consult Arsznieffecting a replacement of the diazo- group by a | CALS, ORGANIC, and ANTIMONIALS, ORGANIC. nitro-radical. A thin aqueous paste of the boro fluorida was added to a auspension of copper in aqueous sodium nitrite. The following reaction | acids (nitrates or sulphates) which are more takes place at room temperature.

In this way a 64% yield of p dinitrobenzene was obtained from p nitraniline, and ethyl p-ammobenzoate furnished a 50% yield of p natrobenzoic forthwith into chloro-pyridines or quinolmes acid (Starkey, J. Amer. Chem. Soc. 1937, 59, 9 Ammophenanthridine (I) does not diazotise 1179). The introduction of a nitron-group is normally in acrd solution, but reacts along the effected by treating a charonum chloride with atth nitrons such, group phenanthridom it alkaline potassium ferricapanide (Hambergee (Morgan and Walls, J.C.S. 1032, 2227). The

salt results in the regeneration of the original amine (Mai, Ber. 1892, 25, 372; 1902, 35, 2593, 3920).

7. Formation of Azolmides.

The introduction of the azimino group N. into the aromatic nucleus is effected by the following reactions: (1) action of hydroxylamine on diazonium aulphates (Mai, Ber. 1892. 25, 372; 1893, 28, 1271; Forster and Fierz. JCS. 1907, 91, 855, 1350); (2) interaction of ammonia and diazonium perbromides; (3) direct action of aodium azide on diazonium salts in excess of sulphurse acid (Nolting and Michael, Ber. 1893, 28, 66 ; Forster and Fierz, J.C S. 1907, 91. 1942). The last of these processes is carried out so readily that the production of an aziminocompound may be used to estimate the number of diazo: groups in a diazonium salt (Meldola and Hawkins, Chem. Soc Proc 1892, 8, 133; Morgan and Micklethwait, J C S 1910, 97, 2560). Aromatic azomides are also produced by the interaction of chloroamine, NH₂Cl, and diazonium chlorides (Forater, J.C.S. 1915, 107, 260), and by removing water from the nitrosoerylhydroxylammes These aryl azomides can be employed syn-

thetically as a means of transforming an amine into the corresponding aminophenol as when warmed with moderately concentrated sulphurio acid, two-thirds of the nitrogen is eliminated and a hydroxyl group is introduced into the nucleus (Friedlander and Zeitlim, Ber, 1894, 27, 192 , Morgan and Porter, J.C S 1915, 107,

The interaction of sodium azide and diazocompound leads directly to the formation of a triazo- derivative in practically quantitative yield. A noteworthy alternative arises when a diazonium salt interacts with an arylsulphonamide in alkaline solution. The reactions are fairly general and the yields favourable.

R SO, NH,+R' N,CI ER SO.H+R'N₃+HCl

R-SO, NH,+2R' N,CI =R SO, N,R'+R'N,+2HCI The former reaction affords a means of pro-

ducing arylaulphinic acids (Dutt. Whitehead

Miscellaneous substitutions, v. J.C.S. 1888, 53,

The primary amines of the triazole and thiazole aeries yield diazonium salts with ovyatable than the corresponding chlorides. Primary Ar N₂ BF₄+NaNO₂=Ar NO₂+N₁+NaBF₄ amines of the pyridine and quinoline series having the azimo group ortho or para with respect to tertiary nitrogen atom yield very unstable diazonium chloridea which decompose other amino- derivatives of phenanthridine are unmethylated dyes, the next three give a readily diazotisable. | mixture of approximately equal parts of un-

Condensations effected by Means of Aromatic Diazonium Salts.

Azo- compounds are formed, to a slight extent, during the Sandmeyer reaction, and this condensation takes place to a greater extent when the usual order of mixing is reversed and the cuprous chloride added to the cold solution of diazonium salt. Aniline, o-chloraniline, and o and p-toluidine give considerable quantities of azo- derivatives, but the nitranilines and their homologues furnish diphenyl compounds (Niementowski, Ber. 1901, 34, 3325; Ullmann and Forgan, ibid. 3802; D.R.P. 126961). Cuprous oxide dissolved in ammonia or hydroxylamine also promotes the diphenyl condensation (Annalen, 1902, 320, 122), diphenyl itself being readily prepared by adding copper, ziuc or iron powder to benzenediazonium sulphate in dilute alcohol (Ber. 1890, 23, 1226); $\beta\beta$ -dinaphthyl is similarly produced from β -diazonaphthalene sulphate (J.C.S. 1895, 67, 653). 2:2'-Dimethyl-1:1'-dianthraquinonyl results from the condensation of 2 mols. of 2-methylanthraquinone-1diazonium sulphate when stirred with acetic anhydrido and copper bronze powder (Scholl and Kunz, Ber. 1907, 40, 1697; B.P. 14578 of 1905).

Coupling of Diazonium Salts.

2:4:6-Trinitrobenzenediazonium salt is highly reactive and couples with mesitylene to form the azo- compound $C_6H_2(NO_2)_3\cdot N_2\cdot C_6H_2Me_3$. This reaction recalls a similar interaction between negatively substituted diazonium salts and certain unsaturated hydrocarbons containing conjugated ethylene linkings. Thus the azo-compound $CH_2\cdot CMe\cdot CMe\cdot CH\cdot N_2\cdot C_6H_4\cdot NO_2$ is readily obtained from dimethylbutadiene and p-nitrobenzenediazonium chloride in glacial acctic acid. This condensation product is reduced by tin and hydrochloric acid to the aminohydrazine,

CH₂:CMe·CMe:CH·NH·NH·C₆H₄·NH₂.

Other diolefins such as butadiene (erythrene), isoprene, and a-methylbutadiene (piperylene) also couple with the active diazo-derivatives from p-nitroaniline and 2:4-dinitroaniline (Meyer and Schoeller, Ber. 1919, 52, [B], 1468).

Diazonium compounds in general couple with ethers of a carboxylated naphthol, thus 2-methoxy-3-naphthoic acid couples in aqueous solution with the diazo-derivative of p-chloro-aniline, 2:4-dichloroaniline, p-nitroaniline, and m-nitroaniline, benzidine, tolidine, o-, and m-nitroanilines, sulphanilic acid, and 5-nitro-o-anisidine. The first six diazo-derivatives give

unmethylated dyes, the next three give a mixture of approximately equal parts of unmethylated and methylated dyes, the last diazo-compound couples without demethylation (Jambuserwala and Mason, J. Soc. Dyers and Col. 1930, 46, 339).

Reactive diazonium salts couple in the side chains of unsaturated aromatic compounds. p-Nitrobenzenediazonium sulphate in alcoholic solution reacts with anethole,

CH3·CH:CH·C6H4·OMe.

to give anisaldehyde-p-nitrophenylhydrazone. With isosafrole, $CH_2O_2:C_6H_3\cdot CH:CH:CH_3$, this diazo- compound gives piperonal-p-nitrophenylhydrazone, and with isoapiole,

CH2O2:C6H(OMe)2·CH:CH·CH3,

apiolaldehydo-p-nitrophenylhydrazone is obtained. A similar condensation occurs with isoeugenol, which yields vanillin-p-nitrophenylhydrazone. These condensations distinguish aromatic compounds containing the propenyl group —CH:CH:CH₃ from those (e.g. safrole) with the allyl side chain —CH₂·CH:CH₂. In the former cases the reactive diazo-derivative, probably in its tautomeric form,

NO₂·C₆H₄·NH·NO,

attacks at the unsaturated linking, forming a hydrazone of the aromatic aldehyde with rupture of the side chain and elimination of acetaldehyde (Quilico and others, Gazzetta, 1928, 58, 389; 1929, 59, 39).

The rate of coupling of diazonium salts with

The rate of coupling of diazonium salts with phenols in buffer solutions of constant ionio strength (μ =0·24) has been measured colorimetrically. Over a wido range of concentration the rate of reaction corresponds with that calculated by means of the bimolecular equation. Rate of coupling is a simple function of the $p_{\rm H}$ of the solution: $\log k = \log k_0 + p_{\rm H}$ (Conant and Petersen, J. Amer. Chem. Soc. 1930, 52, 1220).

Benzenediazonium salts couple with 2:6-diaminopyridine first to a diazoamino-compound, the hydrochloride of which on boiling in aqueous solution gives 2:6-diamino-3(or 4)-benzeneazopyridine (U.S.P. 1820483). 6-Chloro-3-aminopyridine (or the corresponding 6-iodobase) is diazotised and coupled with a variety of reactive pyridine bases such as 2:6-diaminopyridine and its 3-bromo- and 3-iodo- derivatives, to give heterocyclic azo- derivatives which are useful bactericides (B.P. 341598).

The product obtained by tetrazotising ophenylenediamine in nitrosylsulphuric-phosphoric acid solution couples in this strongly acid medium with phenol, β -naphthol, anisole, phenetole and even mesitylene to yield o-diazoazo-compounds so that in such circumstances the bisdiazotised compound should be represented by the formula

in order to explain the one sided coupling, [23993 of 1902; 16995, 27372, of 1903; 4997 of When the acidity of the solution is reduced then a second coupling with a phenolic compound takes place giving riss to o-diazo dyestuffs of outstanding stability towards alkalı (Schoutissen, Rec. Trav. Chim. 1938, 57, 713).

INTERCHANGE OF GROUPS IN DIAZONIUM SALTS

The diazonium salts of the more highly aub atituted amines frequently undergo remarkable transformations, some of which have been suggested for use in the technical production of

azo- compounds. The dinitroanisidines, containing nitro- groups either in ortho- or para- positions with regard to the amino group, exchange these radicals for hydroxyl during the process of diazotisation In certain instances, the methoxyl group present in these compounds undergoes dementary actions that the production of internal diazon better with the production of internal diazon of potassum thiocyanate is converted jate oxides (quinonediazades). The intro-group of potassum thiocyanate is converted jate oxide quinonediazades and which con. 2.4 6 trithrogrambeneediazadum throcyanate tunes the diszoitation process on further and many brommated diszonium oblondle pass quantities of the base (McIdela and others, into chlorusated diszonium bromides (Ber. 1871, J.C.S. 1900, 77, 1172, 1901, 79, 1076, 1902, 81, 130, 233; 1898, 31, 1253; 1900, 33, 505; 67, 988; 1905, 87, 1205). Methoxyl or chlorne J.C.S. 1902, 81, 1376). The last of these 988. 1905, 87, 1200). Methoryl or chlorne J CS. 1902, 81, 1370). De mar ot uner radicals in pare positions with respect to the diazo complex evendered labie by contiguous intro-groups so that gared dance ordees are pro-duced (Klemone, Ber. 1914, 47, 1407; Morgan and Porter, J.CS. 1915, 107, 683), but an access and Porter, J.CS. 1915, 107, 683, but an access unlation of acidic (intro-) groups favours the removal of adjacent nitro groups and the formation of ortho diazo. oxides (Meldola and Reverdin, J C.S. 1910, 67, 1204) A sulphomo group affected, is replaced by hydroxyl in the diszotisation (u) Th of m phenylenediaminedisulphonic acid (B.P. 18283 of 1903), and certain of the dimitronaphthylamines, when diszotised, yield internel diazo- oxidea, owing to the initial replacement of a nitro group by bydroxyl (Gaess and Ammelburg, Ber. 1894, 27, 2211; Friedlander, 1895, 28, 1951; cf. Meldola and Streatfelld, J.C.S. 1895, 67, 998).

The replacement of an acidio (electronegative) substituent by hydroxyl frequently occurs on treating the diazonium salt with transformed in ethyl alcohol; 2 4 6 tribromoalkalts.

$$\begin{array}{c} \text{CIN}_{\bullet} \\ \text{SO}_{\bullet} \\ \text{H} \end{array} \longrightarrow \begin{array}{c} \text{CIN}_{\bullet} \\ \text{SO}_{\bullet} \\ \text{H} \end{array} \longrightarrow \begin{array}{c} \text{OH} \\ \text{OH} \\ \text{SO}_{\bullet} \\ \text{H} \end{array}$$

A sulphonic ecid group may be similarly climinated as in β-naphthylamine-1:5.7 transiphonic acid which, when diazotised and treated with weak alkah, yields naphthylene-2-diazo-1oxide 5.7 disulphonic ocid (G.P. B P. 16811, 20551 of 1901; 1561, 6615, chlorie and, borofluoric acid, notassium tetroxa-

1904). These replacements sometimes take place

even when the diazo- solutions are diluted with water, or spontaneously when the diazonium base is present as the salt of a weak acid (acetate, oxalate, earbonate, etc.) (v. Orton, Proc. Roy. Soc. 1902, 71, 153; J.C.S. 1903, 83, 796; 1907, 91, 1554; Noelting and Battegay, Ber. 1908, 39.

p Chlorobenzenediazoninm thiocyanate

CI C.H.N.CNS undergoes a remarkable molecular transformation when dissolved in alcohol containing a trace of hydrochloric end. The thiocyanste and chlorine radicals change places, and on adding other to the solution, p throeyanobenzene diazonium chlorido is precipitated (Hantzsch, Ber. 1896, 29, 947). Similarly, 2.4 6-tribromo.

ortho- positions being most readily removed. A bromine atom in the meta- position is not

(u) The ease of transformation increases with the number of bromme atoms present.

(us) The transformation constant, calculated from the equation for a unimolocular reaction k=1/t(log A/A-x), increases with the tempera-ture and is also influenced by the solvent, having its minimum value in water, and becoming greater as the aeries of alcohols is ascended.

(iv) Diazonium salts containing two bromine etoms are stable when dry, but are rapidly benzenediazonium chloride becomes transformed in the dry atate,

These changes do not occur either with iodinated diazonium chloridea or with brominated diazonium fluorides (Hantzsch, Ber. 1903,

36, 2069). STABILISED DIAZO. COMPOUNDS,-The technical utdity of stable diazo- compounds has led to many devices for preventing the decomposition of the diazonium group which otherwise occurs on keeping the salts even at the ordinary temperature. When mixed with an equal weight of crystalline alum or aluminium sulphate about two thirds dehydrated, diazo. compounds may he atored for long perioda without deterioration. The diazo derivative of the technically important 5-mitro-o toluidine is atabilised by precipitation as borefluoride (B.P. 316691).

Diazonium borefluorides are rendered more

atable by adding to the solid salt or to its 141750; solution an acid or an acid salt such sa hydrolate, sodium hydrogen sulphate, or dibutyl- acid behave in the peculiar manner indicated in naphthalene- β -sulphonic acid (B.P. 317355). the following diagram: Stable diazonium metallofluorides are obtained by interaction of diazo- solutions with titanofluoric acid or the corresponding complex fluoro acids containing aluminium, stannic, antimonious, antimonic, or zinc fluorides. products are isolated from concentrated solutions by strong cooling (B.P. 332227). The solubility of certain diazonium borofluorides is increased by addition of salts of metals of periodio groups I and III, such as alkali chlorides (including ammonium chloride), alum, or aluminium chloride, B.P. 332630. p-Chlorohenzenesulphonic acid (or its salts) furnishes a stabilising anion when added to diazo- solutions derived from nitranilines, chloronitranilines, chloro-or nitrotoluidines, or any non-sulphonated derivative of these or similar hases. The products are crystalline, water-soluble, stable diazonium p-chlorohenzenesulphonates which may he dried at moderate temperatures (B.P. 265985, 273352). Aromatic diazo-compounds react with aldehyde-hisulphite derivatives of ammonia or amines to furnish soluble stable diazoamino- compounds of the general type

Ar·N:N·NR·CHR'·O·SO,Na.

These diazoamines regenerate diazo-salts by the action of mineral acids. Diazotised 5-nitro-otoluidine couples with the formaldehyde-bisulphite compound of ammonia, methylamine, or dimethylamine. Such diazoamines are used in

dyeing and in the protection of plants from insect pests (B.P. 309610).

A preparation which will liberate a diazocompound when dissolved in cold water is obtained on adding solid sodium nitrite to a mixture containing 1 mol. of an aromatic nitrobase (such as p-nitraniline) rather more than 1 mol. of sulphuric acid (or the equivalent proportions of the sulphate of the base and bisulphates) and a suitable diluent such as partly dehydrated aluminium sulphate or alum, arylsulphonates or mixtures of arylsulphonates and partly dehydrated alum. Such preparations are very stable, and diazotisation is effected on addition of water (G.P. 426033).

Moderately soluhle diazonium fluorosulphonates are obtained by interaction of diazo-compounds and fluorosulphonic acid. These products are insensitive to shock and decompose on heating without explosion; they are used in dyeing and against insect pests (B.P. 303527).

In the manufacture of soluble azo-dyes, diazonium salts are generally employed in aqueous solution, hut solid diazo- compounds, in dry or paste form, have heen introduced to enable dyers to prepare "ingrain" azo- colours which are developed on the textile fibres.

Aryl sulphonic acids have heen employed in this stabilisation of diazo- process, including 1-diazo-2-oxynaphthalene-4-sulphonic and β -naphthalenesulphonic acids naphthalene-1:5-disulphonio acid (Marriott, J. Soc. Dyers and Col. 1936, 52, 172). Anisole-p-

The first change is effected under the influence of alkali when the hemiquinonoid salt dissolves and the solution when acidified yields a precipitate of the insoluble azo-β-naphthol, the sulphonic group having heen eliminated as hisulphite. If the Ar radical contains a nitrogroup, as in the nitranilines, and if excess of alkali is used a deep-seated change occurs and a dihydrophthalazine is formed (Rowe and others, J.C.S. 1931, 1965; 1933, 1067; 1935,

Dried mixtures of an aromatic primary amine, sodium nitrite and an acid salt give rise to diazocompounds on dissolving in water. A mixture of nitrite, β -naphthol, and arylamine-formaldohydebisulphite Ar NH·CH₂·SO₃Na, give a clear solution in weak alkali, but when printed may be developed by acid treatment, which hydrolyses away tho ω -sulphonate group, the primary base, Ar NH_2 , thus liberated is diazotised in acid solution and couples with the naphthol. I.C.I. (B.P. 374497).

LIGHT-SENSITIVE DIAZO- COMPOUNDS.—Tho sensitivity to light of many diazo- compounds has led to various suggestions regarding the application of these compounds to photography.

Diazosulphonates, made from diazonium salts and sodium sulphite, are incapable of coupling in alkaline solutions with phenols or aromatic diamines, but exposure to light removes this inhihition so that a coloured image is produced wherever light has acted. Feer, in 1889, suggested for this purpose an alkaline mixture of ditolylbisdiazosulphonate (derived from tolidine) and resorcinol, a-naphthol, or m-phenylene-diamine. In 1890, Green, Cross and Bevan patented the primuline process hased on the decomposition by light of the diazo- compound of primulin or other thioamines. Paper or linen impregnated with diazoprimuline or diazodehydrothio-p-toluidine was exposed to light, and subsequently immersed in a hath of a phenol or other coupling base. Resorcinol gives an orange tint, β-naphthol a red, 2:4-tolylenediamine a reddish-brown, and α -naphthylamine a violet tint. Of other light-sensitive diazo-compounds may he mentioned o-diazosalicylic acid, Ozalide papers, which have largely displaced blueprint paper, are hased on the principle that a light-sensitive diazoanhydride may be mixed sulphonic acid and acylaminohenzenesulphonic with a phenol or aromatic amine without coupacid also give rise to stable diazonium salts. ling until the mixture is rendered slightly The diazonium salts of β -naphthol- α -sulphonic alkaline. On exposure to light the sensitive diazoanhydride is decomposed except where it is | tained in the hydrolysis of ethyl scetate by protected. The paper hase is then rendered benzened azonium hydroxide, shows that in alkaline by gaseous ammonia when coupling occurs between the undecomposed diazoanhydride and the phenol or aromatic amine with formstion of a coloured azo- compound.

Stabilised disconium salts, such as metallic double salts and borofinorides (see preceding section) from diszotised p-aminoalkylanilmes and p-sminodiphenylamine and its derivatives are employed in preparing diazolype paper. On exposure to hight under a stencil the diazoninm salt is destroyed in parts and the unchanged salt is coupled with phloroglucinol to obtain deep hydroxide to a solution of benzenediazonium shades approaching black or with resorcinel to give brown tints (B P. 281604, 234818, 443995; Chem. Ago, 1936, 34, 198).

Other diazotype papers contain both diazocompound and coupling component The diazoderivative is selected as one giving a diszo- oxide which in acid media is incapable of coupling with a phenol 1. Diazo-2 naphthol 4-sulphonic s cid is an example of this group, it remains uncoupled with a phenol, especially if borne acid or a titanium salt is present Exposed under a tracing the diszo- group is destroyed, and on bringing the paper into a warm ammoniscal atmosphere coupling occurs without eddition of further resgonts (BP 210826, GP 419978). Prints are also obtainable with a diazo com ponent alone, providing that this substance can coupls with its decomposition product. The foregoing 1 diszo 2 rephthol sulphonic acid will do this, as will also diszotised o ammosalicylic acid (G P. 111416) Diazo compounds have also heen utilised in colour photography (Murray, Phot J. 1933, April, and B.P. 363616).

CONSTITUTION OF ARCHATIC DIAZOFTUM SALTS

Atomatic diazonium salta are generally soluble in water to a neutral solution, thus showing, hy an absence of hydrolytic dissociation, that the diszonium base is a stronger base than the aromatic amine from which it was. The benzenoid diszonium salts with colour derived. The strength of the diszonium base less amons (nitrates, sulphates, chlorides) are has been confirmed by physico chemical measurements.

Determinations of the electrical conductivity of solutions of benzenediazonium chiorate and i nitrate show that the benzenediazomum radical is strictly comparable with other qusternary ammonium ions. The rate of migration of the benzenediazonium ion st 25° is 457, the explosibility are considered by Hantzsch to corresponding constants for the methylpyndmium and tetrsmethylammonium ions being substance being regarded as an equilibrium 443 and 436 respectively. The molecular mixture of diazonium salt and syndiazoelectrical conductivity of solutions of diazonium | compound (v. anfra) This equilibrium is dissalts increases with dilution, just as in the case | turbed in the sense of forming more of the colourof the corresponding potassium and ammonium less diszonium salt by lowering the temperature

conductivity experiments, with the results oh- temperature of liquid sir. Apart from their

N/128 solutions at 0°, approximately 33% of the base exists in the ionised condition (Ber. 1890, 23, 3220; 1895, 28, 1737; 1898, 31,

Benzenediazonium (Diazobenzene) behaves

hke an alkalı or a complex ammonium estion Benzenediazonium hydroxide (Diazobenzene hudrate or hydroxide) is only known in solution, and is produced by treating squeous benzepedistonium chloride with the calculated amount of silver oxide, or hy adding barren aulphate (Hantzsch, Ber. 1898, 31, 340). The solution is strongly alkaline, and gives at once an age compound with β-naphthol; the dissolved base decomposes, however, fairly rapidly even

The striking similarity in the physical and chemical properties of the diazonium and quaternary ammonium salts auggests the existence of quinquevalent nitrogen in the former compounds, and on these grounds the following formuls, formerly proposed by Blom-strand (1869), Strecker (1871), and Erlenmeyer (1874), has now been generally adopted for benzenediazonium chloride and its homologues;

although, for many years, the alternative configuration, C. H. N = N Cl, suggested by Kekult, was usually employed.

The close analogy sxisting between the diazonium and alkali selts is shown further b the facility with which both series yield double salts with cobaltio mitrite and with the chlorides of platinum, gold and mercury, Diazonium perhalides are also known, corresponding with the perhalides of potassium, rubidium, cosium and quaternary ammonium bases (Hantzech, Ber. 1895, 28, 1736, 2754).

generally colouriess, but the corresponding bromides, iodides and thiocyanates are coloured, and the deeper the colour the more explosive the salt. When cooled in hound sir, these coloured salts become less roloured; moreover, their aqueous solutions are colourless

The development of colour and increase in indicate a change in constitution, the coloured on the conceptuating possission and ammonium process of another the desiration of the compounds. The affinity constant of henzeodszonium fluenzach, Ber. 1897, 33, 2179; 1900, colour, these salts have all the properties of the simpler diazonium salts (neutral solutions, coupling with phenols, etc.). These facts indicate that increase in the complexity of the organic portion of the diazonium cation leads to the development of colour in normal diazonium salts.

STEREOCHEMISTRY OF DIAZO- DERIVATIVES .-The stereochemistry of diazo- compounds advocated by Hantzsch, and accepted generally by workers in this field of organic chemistry, is based on the existence of two isomeric forms of aromatic diazocyanides, diazosulphonates and diazo-oxides. Of these two series the more stable anti-diazo- compounds have frequently been prepared in a state of purity so that their existence has been confirmed by many analyses. The labile syn-diazo- compounds, on the other hand, are much less readily isolated, and analytical data in support of their existence as definite chemical entities are not always obtainable. 4'-Amino-1-phenyl-5-methylbenzothiazole (dehydrothio-p-toluidine) was shown by its discoverer A. G. Green (1889) to give rise to a diazo-chloride quite stable in solution at the ordinary temperature.

When suspended in absolute alcohol and diazotised with ethyl nitrite, the hydrochloride of the foregoing aminothiazole yields two isomeric compounds, an insoluble diazochloride and a soluble one precipitated by ether. Synl-phenyl-5-methylbenzothiazole-4'-diazochloride decomposes at 139°, dissolves readily in water, couples immediately with alkaline a-naphthol, and evolves nitrogen with copper bronze, giving 4'-chloro-I-phenyl-5-methylbenzothiazole. Antil-phenyl-5-methylbenzothiazole-4'-diazochloride decomposes at 270°, is insoluble in water, does not couple with alkaline β -naphthol until after treatment with hydrochloric acid. With copper bronze and this acid it remained unchanged for several hours, but after a week yielded the chloro-derivative.

Ice-cold solutions of the syn-diazochloride when treated with sodium sulphite gave a sodium syn-1-phenyl-5-methylbenzothiazole-4'-diazosulphonate, a salt which couples with β -naphthol. At the ordinary temperatures (about 20°) a non-coupling anti-diazosulphonate is obtained. Similar experimental conditions lead to the formation of syn- and anti-1-phenyl-5-methylbenzothiazole-4'-diazocyanides. The syn-diazocyanide is a brick-red product, very soluble in water and decomposes at 131°, it couples readily with alkaline β -naphthol and readily loses nitrogen with copper bronze. The anti-diazocyanide, a deep orange precipitate insoluble in water, melts at 175°, does not couple with alkaline β -naphthol, and evolves nitrogen only very slowly with copper bronze. Accepting the stereochemical hypothesis these three pairs of syn- and anti-diazo- compounds may be thus represented:

The solubility in water of the syn-diazochloride and -diazocyanide suggests that these compounds pass readily into the diazonium condition, D being the thiazole radical (Morgan and Webster, J.C.S. 1921, 119, 1071).

$$\begin{array}{c|c}
D-N \\
\parallel xH_2O
\end{array} \rightleftharpoons \begin{bmatrix}
D-N \\
\parallel xH_2O
\end{bmatrix} CI$$

In this benzothiazole series there was also qualitative evidence of the existence of syn- and anti-diazo-oxides (diazotates). For the constitution of diazo-oxides (diazotates), see Angeli, Ber. 1926, 59 [B], 1400; Hantzsch, ibid. 1929, 62, [B], 1235; Quilico, Gazzetta, 1931, 61, 326.

(Coloured diazonium salts, v. Hantzsch, Ber. 1900, 33, 3183; 1901, 34, 4168; Annalen, 1900, 312, 126; Cain, J.C.S. 1905, 87, 5; Morgan and others, ibid. 1907, 91, 1311, 1505; 1908, 93, 614; 1909, 95, 1319.)

The idea embodied in Blomstrand's diazonium formula (v. supra) fails to express one important point, namely, the dependence of the stability of the diazo- complex on the presence of an unsaturated group. Yet this is a matter of fundamental importance, for hitherto no amino in which the basic nitrogen is attached to a fully saturated ring or complex has yielded a diazonium salt.

Cain has suggested an alternative formula, (2), giving expression to this circumstance (J.C.S. 1907, 91, 1051), in which the diazonium salt is represented as having a para-hemiquinonoid constitution; and this conception has been extended by Morgan (*ibid.* 1908, 93, 617; 1910, 97, 2563) so as to include the equally admissible ortho-hemiquinonoid configurations (1) and (3):

Euler has also advocated a para-hemiquinonoid structure for diazonium salts (Ber. 1908, 41, 3979).

Non-aromatic Diazonium Salts.

The foregoing hypotheses of the constitution of aromatic diazonium salts, which bases their existence on the unsaturated nature of the aromatic complex, accord with the fact that certain non-aromatic primary amines containing unsaturated organic groups manifest, in some degree, the property of diazotisability. Among

IV.

these hases are aminotriazole (I) and its deriva-

(Thiele and Manchot, Annalen, 1898, 203, 32; Knorr and Stok, Joh. 1890, 233, 67; cf Ber 1900, 33, 1158; 1900, 39, 2925; Annalen, 1900, 131, 133; In addition to amon-denvatives of the pyraxolme group (Michaelis, Annalen, 1906, 300, 288), the 4 and 5 amin derivatives of the pyraxole group gree rise to remarkably stable duancinium saits (Morgan and Redly), 258 1914, 105, 435; Michaelis and others, Annalen, 1913, 537, 1; 1915, 407, 229, 274; 1904, J. pr. Chem., 1914, [61, 80, 500). 2-Ammedianch, and present the range of the property of the

1231:

Pyrazole Series.—Among pyrazole bases the effect of orientation on the etablity of the diazonium salts is well marked.

When diarotised in the usual manner, 4 minos-3.6-dimethylpyratole (I) yelds the diazonam chloride (II) stable in either hot aqueous solutions or up to 100° in the dyr state. When the diaro-group is in poetion 5 the product is much less stable, for 1-phenyl-3 methyl-4-ethylpyratole-5-diazonium chloride decomposes quickly stem at poom temperature. If the 4-ethyl group is replaced by labile hydrogen as in 1-phenyl 3-methyl-5-aminopyrazole bydrochloride (III), then introns and yields only 12% of diazonium sait (IV), the man product (Son of diazonium sait (IV), the man product (Son

being the imino-oxime (V) (Mohr, J. pr. Chem. 1914, 90, (si), 509).

ino-Drainle Striet.—3.5 Dimethylaronanole disnosum-5-sulphonate, has been converted into 4 nitro-3.5 dimethylaronanole, and the latter reduced by amalgamated alumnum ond most ether to 4 amuno-3.5-dimethyliaronanole (II), which diarotises to the very soluble diaronum chloride (III). This diarotise that the very soluble diaronum chloride (III). This diarotise that the summan of the very soluble diaronum chloride.

3 5-Dimethyl iso-oxazolo-4-diazonium chloride interacta readily with sodium axide or potassium totale to yield respectively 4-triazo-3-5-dimethylico-oxazolo (III) and 4 ido-3-5-dimethylico-oxazolo (III) and 4 ido-3-5-dimethylico-oxazolo (III) and 4 ido-3-5-dimethylico-oxazolo (III) and 4 ido-3-5-dimethylico-oxazolo (III) and 1 ido-3-dimethylico-oxazolo (III) and 1 ido-3-dimethylico-oxazolo (IV) and 1 ido-3-dimethylico-oxazolo (IV) and 1 ido-3-dimethylico-oxazolo (IV) (Morgan and Burgess, J.C.S. 1921, 1938, 1546).

Datunct evidence of diazofisability has been adduced in the indole and pyrrole series [Angele and d'Angelo, Atti R. Accad. Lines, 1904, [v], 1, 253 s. Angelico, ibid. [105, [v], 14, il. 167]. It has also been shown (K. A. Hofmann, Host and Reth. Ber. 1910, 43, 622, 1087) that ammoguanidine given rise to distornium salts, derived, bowever, not from the salts of ammoguanidine itself, but from a more compilex molecule containing two guantidine residues, and having a taking two guantidine residues, and having a naturation. The discount natural, for adample, is represented by the formula:

NH C(NH*)·NH—NH N N

containing a diazo- complex associated with an open chain having three unsaturated linkings.

II. Cyclic Diazo. Compounds.

In certain instances, the action of nitrous acid on substituted aromatic animes leads to the formation of compounds having their disconitrogen in a cyclic complex, this result being due to the subsequent interaction of a substatuent group.

1. INTERNAL DIAZONIUM SALTS.

Sulphonated amines of the benzen and maphthalene series furnah diazo derivative which are generally minamed diazo-nighonic aceds. These compounds, however, do not contain nonic hydrogen, and are really internal diazonium sale; No. 1, the so-called diazoniump-nighonate, whits No. 2 is naphthalene-1 diazonium-5-nighonate.

great technical importance in the manufacture of azo- dycs. They are prepared either by adding sodium nitrite to an acid suspension of the amino sulphonic acid or hy acidifying a solution containing nitrite and the alkali salt of the aminosulphonic acid (Rev. prod. chim. 1917,

Of a similar nature are the cyclic diazoderivatives obtained by diazotising aromatic aminocarhoxylic acids (cf. Bülow and Haas, Ber. 1911, 44, 601), e.g. "diazo-anthranilic acid"

or benzenediazonium-o-carboxylate

formula (3 a).

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which is employed in the production of methyl red (v. CHEMICAL ANALYSIS, Acidimetry and Alkalimetry, Vol. II, p. 641).

These internal diazonium salts are now often formulated as polar substances, Formula (1) heing modified as N₂·C₆H₄·SO₃ and the lastmentioned diazonium carhoxylate as shown in

2. CYCLIC DIAZO- COMPOUNDS DUE TO INTERNAL CONDENSATION.

Aromatic ortho- and peri-diamines do not yield diazonium salts except in special circumstances (Schoutissen, J. Amer. Chem. Soc. 1935, 55, 4535; Hodgson and Walker, J.C.S. 1935, 530), inasmuch as their interaction with nitrous acid leads very readily to the formation of cyclic diazoimines (Azimino- compounds)

Phenylene-o-diazoimine (1) (Aziminobenzene) and naphthylene-1:8-diazoimine (2) (1:8-Azimino-naphthalene) are typical members

of this series of diazo- compounds.

The triazole rings thus set up are remarkably stable, and in the case of the ortho- derivatives (1) are not opened hy mineral acids without complete rupture of the molecule. The peridiazoimines (2) occupy a position intermediate between the ortho-diazoimines (1) and the paradiazoimines (v. infra).

The 1:2- and 2:3-naphthylene-diamines furnish respectively 1:2- and 2:3-naphthylene-diazoimines (1:2- and 2:3-azimino-naphthalenes). (Constitution of ortho-diazoimines and their acyl-derivatives, Griess, Ber. 1882, 15, 2195; Kekulé, Lehrhuch, ii, 739; Morgan and Godden, J.C.S. 1910, 97, 2557; cf. Ber. 1874, 7, 316; 1876, 9, 221; 1887, 20, 2999; 1894, 27, 874; Annalen, 1889, 255, 344; J.C.S. 1906, 89, 4). Although Griess's formulation for the azimino- compounds

$$C_6H_4$$
 NH still persists in chemical

These internal diazonium sulphonates are of literature, it has been disproved completely, and that of Kekulé substantiated by the preparation of three pairs of isomerides having respectively the following formulæ, where R=acetyl, benzoyl, or henzenesulphonyl.

This form of isomerism could not exist with Griess's symmetrical configuration (Morgan, Micklethwait and Scharff, J.C.S. 1913, 103, 1394; 1914, 105, 117). Although these diazoimines do not couple with phenols and aromatic bases to form azo- dyes, yet employment has been found for them in the production of synthetic carbazole derivatives which are obtained on heating these cyclic diazo- compounds (Ullmann, Ber. 1898, 31, 1697; 1914, 47, 380).

Similar stable cyclic diazo- compounds are ohtained hy diazotising the ortho-aminothiophenols, o-phenylenediazosulphide being a colourless crystalline substance with a sweetish odour and feebly hasic properties

(Jacobson, Annalen, 1893, 277, 209, 218, 232, **237).**

3. CYCLIC DIAZO. COMPOUNDS DUE TO COMBINED Internal Condensation and Salt Forma-TION.

In this group are found certain diazoderivatives of amines containing slightly acidio substituents in ortho- or para- but not metapositions with respect to the aminic nitrogen.

(i) Internal diazo-oxides are produced from ortho- and para- aminophenols and their derivatives. In some cases an intermediate diazonium salt is formed which condenses to the diazo-oxide on treatment with silver oxide or an alkali (Schmitt, Ber. 1868, 1, 67; Hantzsch and Davidson, *ibid.* 1896, 29, 1522; Cameron, Amer. Chem. J. 1898, 20, 229). The first diazocompound prepared by Griess (Annalen, 1858, 106, 123), namely, dinitrobenzenediazo-oxide,

 $(NO_2)_2C_6H_2$, belongs to this class; it is

produced by diazotising picramic acid, and has been utilised in the production of azo- colours. The 1:2-, 2:1-, 2:3-, and 1:8-amino-naphthols and their sulphonic acids similarly give rise to internal diazo oxides which are so stable that they can be nitrated. These internal diazooxides are readily transformed into diazonium salts by mineral acids, and are therefore available in the production of azo- dyes.

Internal diazo-oxides can also be produced by leaving the diazonium salts of highly suhstituted amines (e.g. 2:4:6-trichloroaniline) in >NH still persists in chemical acid (Orton, Proc. Roy. Soc. 1903, 71, 153;

J.C.S. 1903, 83, 796).

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quinonediazides (11):

(cf. Wolff, Annalen, 1900, 312, 126; Morgan and Micklethwait, J.C.S. 1908, 93, 607).

Repeated attempts to prepare internal diszo-oxides from m-aminophenol and its intro-derivatives, which have been quito unsuccessful, strengthen the view that these diazo- derivatives are to be regarded as quinonediaze exides (II), with the reservation that similar compounds are producible from 1:8 aminonaphthol, 'If acid,' and other peri- derivatives (Morgan and Porter, J.C.S. 1915, 107, 645). In the case of ammo phenoisulphonic sends the NH, OH =1:2 compounds give these coloured e-quinonediazides; the NH₂OH=1:4 compounds yield diazonium aulphonates changing to p-quinonediazides on partial neutralisation; the NH₂ OH=1.3 compounds give rise only to diazonium sulphonates which lose diazo-nitrogen and furnish complex azo. derivatives on treatment with alkalı (Morgan and Tomlins, J.C.S. 1917, 111, 497).

AROMATIO DIAZO-ORIDES. - 2.4 Dinitro cnaphtbylamine is a feeble base which is diazotised by adding nitrosyl sulphate (chamber crystals) or dry sodium nitrate to the base dissolved in cold concentrated sulphuric acid, the agintion being subsequently diluted with frag. The following alternative formula ments of ice. The liquid is then poured into a IV) have been given to this product: large excess of 100-water, when a crystalline brownish yellow precipitate of 4 nitronaphthalens I diazo 2 oxide (1) is obtained,

$$\bigcup_{I_{i}}^{N-N} \bigcup_{NO_{a}}^{N-N} \bigcup_{II_{i}}^{N-N} \bigcup_{N=N}^{N-N} \bigcup_{N=$$

This remarkably stable diszo-oxide couples only with the more reactive phenols such as reservinol phloroglucinol and 1:3 dihydroxynaphthalene, and in these instances coupling occurs not only and in these mataness coupling occurs not only in alkalme adultions, but also, and with great facility, in presence of atrong acids, for example, in glacial acetic acid containing concentrated mineral acid (Morgan and Evens, J.C.S. 1919, 115, 1126). The technically important naphthalenediazo-2-oxide-4-sulphonie acrd II forms an acidic brown copper compound which gives no precipitate with acdium hydroxide and a yellow solution with ammonia (Battegsy and Schmidt, Bull. Soc. chim. 1927, (iv), 41, 205).
For the diazotisation of aminophenols which

are readily oxidised to quinones, nitrites are employed in presence of mineral acids and salts of copper, iron, zinc or similar metals. The resulting diazo solutions may be used either for compling to form azo dyes or for displacement of the diazo group by bydrogen, halogons of other substituents (G.P. 431513).

These diazo derivatives have been formu- | phenylstihine dichlorides diazotise to yield comated either as cyclic diazo-oxides (1) or as pounds of the type ClaX.C.H., N.Cl (where quinonediazides (11):

X=As or Sb). The arsenical derivative on treatment with sodium hydrogen carbonete is converted into a p-phonylarsonic acid

which on reduction with sulphur dioxide in presence of iodine yields p-phenylenechloro-arsine (I; X=AsCt).

3-Amino-4-hydroxyphenylarsinie acid when similarly treated gives first the diago, aalt AsCl₂C₂H₁(OH)N₂Cl, which readily loses hydrogen chloride to pass into the arenical quinonediazide (II) (Sebmidt and Hoffmann, Ber. 1926, 58, [B], 555, 560).
(a) peroDiazolmines and diazolmides,

p-Diazolminobenzene and derivatives .-The earliest known derivative of p-diazoimino. benzeno was obtained by the action of alkali on diazonium anlts of p-aminodiphenylamine (Ikuta, Annalen, 1888, 243, 282; Hantzsch, Ber. 1902,

The following alternative formulæ (III and

corresponding respectively with the foregoing formulæ (I and II) for the internal diazo oxides Successive nitration of the phenyl group gradually increases the stability of these p. diazoimines (Morgan and Micklethwait, J.CS. 1908, 93, 604).

The aryisnipbonyl derivatives,

were first discovered by Morgan and Micklethwart in 1905, and the simplest acyl derivative, formyl-p diazoiminobenzene, was isolated by Morgan and Upton in 1917).

The para-diazoimides are produced from the acyl- and aryisulphonyl- derivatives of the para-diamines, and similar products are obtained from the aryisulphonyl-1:8-naphthylendifrom the arylsulphonyl-1:8-naphthylenedi-amines. The production of these diazomides takes place in two stages, the diazonium salt first produced being decomposed by sodium scetate or mild atkali, yielding the Internal condensation product :

$$R \cdot SO_a \cdot NH \cdot C_aH_a \cdot N_a \cdot C_i \stackrel{NaAc}{\rightleftharpoons} R \cdot SO_a \cdot N \cdot C_aH_aN_a$$
 V , VI .

These para-diazoimides (VI) are yellow, Aminophenylarsino dichlorides and amino- sparingly soluble substances readily reconverted

by strong acids into the corresponding diazonium | 155°. They combine additively with phenols, naphthols, naphthylamines and their sulphonic acids, yielding azo- colouring matters (Morgan and Micklethwait, J. Soc. Dyers and Col. 1909, 25, 107; cf. J.C.S. 1905, 87, 74, 921, 1302; 1906, 89, 1162; 1907, 91, 1509; 1908, 93, 615; Badische Anilin- und Soda-Fabrik. G.P. 205037).

The foregoing general reaction has been extended further by the device of diazotising with liquid nitrogen trioxide in an anhydrous solvent such as acetone. In this way the simplest members of the series Ac.N:C₆H₄:N₂ have been obtained containing formyl, acetyl, and benzoyl groups (Morgan, Upton, and Cleage, J.C.S. 1917, 111, 187; 1918, 113, 588.

3-Aminocarbazole hydrochloride gives rise to a very stable yellow diazonium chloride

C₁₂H₈N₃Cl,3H₂O

which becomes anhydrous when stored in vacuo over sulphuric acid. With sodium azide it yields a colourless 3-triazocarbazole, m.p. 176-177°.

$$\begin{array}{c}
NH \\
I. \\
-HCI \\
+HCI
\end{array}$$

$$\begin{array}{c}
N \\
N \\
\parallel N
\end{array}$$

$$\begin{array}{c}
N \\
\parallel N \\
\parallel N
\end{array}$$

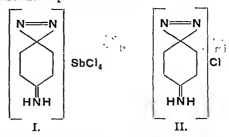
When carbazole-3-diazonium chloride (I) is treated in the cold with ammonia in the absence of light, hydrogen chloride is removed and carhazole-3-diazoimine (II) is precipitated in bright orange-red needles darkening at 80-90° and exploding at 95°. This diazoimine is extremely photo-sensitive and decomposes almost immediately in sunlight. It couples with phenols and reactive aromatic amines, and on addition of hydrochloric acid regenerates the diazonium chloride (I) (Morgan and Read,

J.C.S. 1922, 121, 2711).

In 1926 W. H. Gray obtained the hydrochloride and double antimony chloride of p-diazoiminobenzene itself by the following series of reactions.

The hydrochloride of acetyl-p-phenylenediamine was diazotised and treated with antimony trichloride when bis-p-acetaminobenzenediazonium chloride antimony chloride separated as a yellow crystalline precipitate, m.p. 147°. decomposed with water this double salt furnished p-acetaminobenzenediazonium chloride, 131°, which on hydrolysis with boiling dilute hydrochloric acid yielded p-diazoiminobenzenehydrochloride-antimony trichloride (I), decomwith water furnished p-diazoiminobenzene hydro-

Both the corresponding picrate chromate explode at 160°.



In bright sunlight acetic anhydride converts the foregoing hydrochloride into p-acetaminobenzenediazonium chloride.

Acetyl-m-phenylenediamine hydrochloride when diazotised and combined with antimony trichleride gives bis-m-acetaminobenzenediazonium hydrochloride-antimony trichloride, m.p. 94°, which, unlike its para-isomeride, is decomposed on warming with dilute acid, but when treated successively with cold water and chromic acid yields an explosive m-acetaminobenzene-diazonium chromate. There is no indication in this meta-series of the formation of a diazoimine similar to the above compound (II) (J.C.S. 1926, 3174).

III. Diazoamines.

The diazoamines or diazoamino- compounds have the general formula XN2 NHY, and may be either aromatic, mixed aromatic-aliphatic, or aliphatic in character, according as to whether X and Y are aromatic or aliphatic hydrocarbon radicals.

1. AROMATIC DIAZOAMINES.

The aromatic diazoamines are produced (1) by the action of nitrous acid (1 mol.) on two molecular proportions of a primary benze-

noid amine; (2) by coupling a diazonium salt with a primary or secondary benzenoid amine.

Diazoaminobenzene, C₆H₅·N₂·NH·C₆H₅, yellow crystals, m.p. 98°, is produced in quantitative yield by dissolving 10 parts of aniline in two parts of water containing 12 parts of hydrogen chloride, diazotising with 8 parts of sodium nitrite, and adding the diazo- solution to a solution of 10 parts of aniline in the calculated amount of hydrochleric acid. The diazoamine precipitated on the addition of sodium acetate. (Absorption spectrum of diazoaminobenzene, Purvis, J.C.S. 1914, 105, 590.)

When warmed with aniline containing aniline hydrochloride, diazoaminobenzene is formed into p-aminoazobenzene, the velocity of transformation obeying the law of unimolecular reactions. When the transformation occurs below 40° a small proportion (about 4%) of o-aminoazobenzene is produced (F. H. Witt, Ber. 1912, 45, 2380; 1913, 46, 2557).

Diazoaminobenzene is conveniently prepared in a state of purity by passing carbon dioxide posing at 179°, and this double salt on treatment | into an aqueous alcoholic solution of aniline and sodium nitrite. The product recrystallised from chloride (II), a yellow substance exploding at alkaline alcohol separates in lemon yellow

needles, mp. 995-100°. Diluto acids cause a lowering of this melting point. When the action of cold glacial acetic acid on diszoaminebenzene is allowed to proceed until it is nearly diphenylamine, ortho and para-aminodiphenyls all transformed into aminoszobenzene a non-basic product of this reaction is isolated on diphenyl with evolution of nitrogen. More diluting the liquid with water, when a tarry mass recently azobenzene has been identified among is deposited from which by repeated crystallisation a deep red substance, benzenediazoaminoazobenzene, Callan, C.H., NH.N., C.H., is obtained (m.p. 1195°), identical with the compound produced by coupling diazotised aminoacobenzene and anilme. It is extremely difficult to separato this substance from diazoaminobenzene, and mixtures of the two cubstances which melt lower than either compound have in the past suggested the existence of an isomeric and more fusible diazoaminobenzene (Earl, J. Proc. Roy. Soc., New South Wales, 1930, 64, 97). Interesting observations on the transformation of deazoaminobenzene into aminoazobenzene have also been made by Rosenhauer (Ber. 1928, 61, [B], 392; 1930, 63, [B], 1056; 1931, 64, [B], 1438), who holds that this transformation is due to primary fission into diazonium salt and aniline followed by nuclear coupling and not to isomeric change. The conversion is effected not only by glacial acetic acid, but also by dilute aqueona acetic acid or formso acid, never by quinoline and its hydrochloride. Benzenediazonium chloride and aniline hydrochloride couple at 0° to give aminoszobenzene hydrochloride without intermediate formation of diazoaminobenzene. About 70% of aminoazobenzene is ohtainahle in acetic acid solution if 3-4 mols, of aniline are added.

Diazoamino-p-toluene,

similarly prepared to diszoaminobenzene, undergoes transformation into an ortho-azo- compound hut the velocity of this change is only about one-ninth of that of the preceding transforma-

tion to para azo- derivative.

The coupling of diazonlum salts with meta-phenylenediamine, diphenylamine, the naphtbylammes and their alkyl derivatives leads directly to the production of azo- derivatives; in these cases, intermediate diazoemines have not been isolated, except when the reactive ortho- or para positions are already substituted (Morgan, J.CS. 1902, 81, 91; 1907, 91, 370).

When an aromatic base, XNH, is coupled with a diazonium salt, YN,Cl, the resulting diazonium is identical with that produced from XN,Cl and YNH, and is generally regarded as being an equilibrium mixture of the two iso-merides XNH N₂Y and XN₂ NHY. This ex-planation assumes the migration of the diazogroup which has been observed to occur in coveral instances (Griess, Ber. 1882, 15, 2190 : Schraube and Fritsch, 1896, 29, 287; Hantzsch aud F. M. Perkin, 1897, 30, 1412).

(For the constitution of the unsymmetrical arometic diazoamines, XNH N.Y. v. Meddola and Streatfeld, J.C.S. 1886, 49, 624; 1887, 57, 102, 434; 1886, 53, 664; 1889, 55, 412; 1890, 57, 785; cf. Ferster and Garland, shid. 1909, 95, 2051.)

Pyrolysis of Diazoamino benzene and -p-toluene .- In 1892 Hirsch pyrolysed diszoaminobenzene aud obtained p-ammoazobenzene.

these pyrolytic products.

A quantitative study of the pyrolysis of diagoamino-p toluene dissolved in p toluidine at 160° has shown that the more volatile products are nitrogen, ammonia, and toluene, whereas the less volstile products are azo-p-toluene, pditolylamine, 6-amino-3.6'-dimethylazobenzene and two isomeric aminoditolyls (15% yield). One isomeride obtained in very small amount has not been orientated; the more plentiful isomerade is 4'.5-dimethyl 2-xenylamine, the oily triazo- compound of which is converted pyrogenically 2.6 dimethylcarhazole into (Morgan and Walls, J.C.S. 1930, 1502).

2. ALIPHATIC-AROMATIC DIAZOAMINES.

The diszoamines containing one alighatic and one aromatic group can be produced by (I) coupling a diazonium salt with an aliphatic amine (Goldschmidt and Holm, Ber. 1888, 21, 1016, 1112; Goldsmidt and Badl, 1889, 22, 938; Bamberger and Müller, ibid. 1302); (2) treating an azide with Grignard reagent :

$$RN < \bigcap_{N} + R' \cdot MgBr \rightarrow RN(MgBr) N.NR'$$

$$\xrightarrow{H_2O} RNH \cdot N: NR' + MgBr \cdot OH$$

(Dimroth, Ber. 1903, 36, 909; 1905, 23, 670, 2326; 1907, 40, 2390). Diazoamines present the possibility of stereoisomersm, but their relative stability and the fect that they couple only slowly with \$-naphthol confirm the view that they are generally anti-diazo- compounds.

(For atereosomerio diazoemines v. Orloff, J. Russ. Phys. Chem. Soc. 1906, 58, 578; Vaubel,

Z. angew. Chem. 1900, 13, 762; 1902, 15, 1209.1

Aromatic budiazoamino compounds. (C.H. N.)2N C.H., have been obtained by the action of two molecular proportions of a diazonium ealt on one of an aromatic amme (Hantzsch, Ber. 1894, 27, 1863; von Pechmann and

Frobenius, 1895, 28, 170). Mixed aliphatic-aromatic biediazoamino derivatives (v. Goldschmidt et al., Ber. 1888, 21,

1016; 1889, 22, 933; 1907, 40, 2390). Diazo-4-asmicarbazines (v. Forster, J.C.S. 1906, 89, 223).

3. ALIPHATIC DIAZOAMINES.

Dimroth's general method may be applied to the production of purely aliphatic diazoamino compounde. In this wey, the simplest member of the series, diazoaminomethane (dimethyltriazene), CH₃N:N·NH CH₃, has been prepared from methyl szide and magnesium methyl iodide. This substance, which is isolated by distilling its copper derivative with diazoaminobenzene under reduced pressure, is a colourless liquid, solidifying at -12° and boiling at 92° (Dimroth, Ber. 1905, 38, 1575; 1906, 39, 3905).

IV. Binuclear Diazo-oxides.

When a diazonium salt couples with a phenol the product is generally an ortho- or a parahydroxyazo- compound (the C-azo- derivative), but in all probability, the O-azo- derivative is first produced, for in a few instances, when the velocity of transformation has been lessened by substitution, the intermediate O-azo- compound or diazo-oxide has been isolated. p-Bromobenzenediazonium chloride and p-nitrophenol give p-bromobenzenediazo-4-oxymitrobenzene

which, at 80°, becomes transformed into its isomeride, p-bromobenzene-2-azo-4-nitrophenol

(Dimroth and Hartmann, Ber. 1908, 41, 4027; Auwers, ibid. 4304).

Occasionally traces of these intermediate diazoamines and diazo-oxides can be detected in the commercial azo- colouring matters (Vaubel, Z. Farben. Textil. Ind. 1902, 1, 3).

V. Metallic Derivatives of Diazo-Compounds.

In 1894 Schraube and Schmidt (Ber. 1894, 27, 514) found that on adding a 10% solution of p-nutropenzenediazonium chloride to 18% aqueous sodium hydroxide at 50-60°, a sodium diazo- derivativa constant at 50-60°, a sodium diazo- derivative separated in yellow bronzy leaflets, to which they gave the formula

regarding the compound as the sodium derivative of p-nitrophenylnitrosamine. This view of the constitution of the product was confirmed by the action on the compound of methyl iodide, when p-nitrophenylmethylnitrosamine,

$$NO_2 \cdot C_6 H_4 \cdot N(CH_3) \cdot NO_7$$

was obtained.

This sodium salt, which was regarded by Hantzsch as having the constitution of an antidiazo- compound, sodium p-nitrobenzenediazo-oxide (v. infra), has been manufactured for use in the production of "nitrosamine red" (Badische Anilin- und Soda-Fabrik, G.P. 78874, 80263, 81134, 81202). The addition of excess of mineral acid to this metallic diazo- derivative determines the regeneration of the original diazonium salt, but treatment with dilute acetic acid leads to the formation of a yellow product, p-nitrophenylnitrosamine, NO₂·C₆H₄·NH·NO, a substance showing little tendency to couple with β-naphthol.

The silver derivative obtained from the foregoing sodium salt yields on methylation an oxygen ether NO2·C6H4·N:N·O·CH3, a reaction suggesting the possibility of tautomeric change. In many instances two isomeric modifications of these metallic diazo- derivatives have been isolated.

Benzenediazonium chloride and cold concentrated potassium hydroxide yield the normal labile form of potassium benzenediazo-oxide (I), which readily couples with β-naphthol. The stable modification of potassium benzenediazo-oxide (II) is produced by heating the strongly alkaline solution of the normal or labile salt at 130-140°; this isomeride couples only very slowly with β -naphthol.

According to Hantzsch, these isomerides have respectively the following formulæ:

I. Syn- (labile form).

II. Anti- (stable form).

Dobbie and Tinkler (J.C.S. 1905, 87, 273) have found, however, that these isomeric potassium salts have totally different ultra-violet absorption spectra. But as stereo-isomerides, like the benzaldoximes, have identical absorption spectra, it seems probable that the above isomeric salts are structurally dissimilar. Confirmation of the stereochemical theory would be obtained by isolating the anti-diazohydroxide from the stable (anti-) potassium salt, but on treating this with an acid, the nitrosamine separates. Moreover, the product supposed by Hantzsch and Pohl to be an anti-diazohydroxide (Ber. 1902, 35, 2964) was shown by Orton to be a mixture of quinonediazide and a hydroxyazo- compound (Proc. Roy. Soc. 1903, 71, 153; J.C.S. 1903, 83, Ż96).

The existence of the syn-diazohydroxides is doubtful, and diazonium hydroxides (v. supra) are only known in solution

Diazo-anhydrides, RN:N·O·N:NR RN-O·N:NR

, are very unstable explosive com-Ν pounds produced by the action of acetic acid on the metallic syn-diazo-oxides. The anti-diazooxides, when thus treated, give nitrosamines.

Aromatic diazoic acids, Ar·N₂O₂H. Benzenediazoic acid, produced by oxidising potassium benzenesyndiazo-oxide or isodiazo-oxide with potassium ferricyanide in alkaline solution, is obtained in sparingly soluble white leaflets (m.p. 46°) dissolving readily in organic solvents or alkalis. With mineral acids it is transformed into o- and p-nitroaniline, and with hypochlorites it yields 4-chloro-2-nitroaniline. Its sodium salt gives the α - or N-ester, C_6H_5 -N(CH₃)·NO₂, the silver salt furnishes the β - or O-ester, C_6H_6 -N:NOO-CH₃. Since the 4-chloro-2-nitroaniline. acid is also obtained by the action of nitrogen pentoxide on aniline it is probably a tautomeric substance:

$$C_6H_5\cdot NH\cdot NO_2\rightleftharpoons C_6H_5\cdot N:NO_9H$$

(Bamberger, Ber. 1893, 26, 471; 1894, 27, 359 914, 584, 1273; Hantzsch, Ber. 1902, 35, 258).

VI. Diazocyanides.

The addition of the calculated amount of potassium oyanide to a cold acid solution of a diazonium salt determines the formation of a sparingly soluble diazocyanide. In instances, two modifications of the diazocyanides can be distinguished, and these varieties | diazonyanides is attructure), the sym-compounds are regarded by Hautzch as being stereines—finely teoryanides R-N:N:NC and their smir-incris forms. p Chlorobenzenedasonium chlor-isomerides, cyanides R-N:N CN (R-chamide, ide, from p-chloranilme, yields below -5 the Ber. 1995, 23, 801; Orton, J.C.S. 1903, 83, yellow | lable | sym-p-chlorobenzenedazo-1805|. cyanide (1), which readily evolves nitrogen,

vields p-chlorobenzonitirale on treatment with copper powder, and readily passes into the stable anti-p-chlorobenzenedlazocyanide (11). The latter substance, which is not affected by copper powder, may be distilled in steam without decomposition (Hantzsch and Schulze, Ber. 1895, 28, 666).

Confirmation of this stereochemical theory of the constitution of the diazocyanides has been obtained by a study of the cyanides derived from p methoxybenzenediazonium ehloride. This salt with potassium cyanide in alcoholic solution yields the syn diazocyanide (III), an orange-red insoluble substance (m.p. 51°), which couples with 8 naphthol and slowly changes into the non coupling anti-diagooyanide (IV), a brownish red compound, melting at 121°.

The existence of a third isomeric cyanide is indicated by evaporating at the ordinary temperature in the presence of excess of hydrocyanio acid, an aqueous solution of p-methoxy-benzenediazonium hydroxide. The colourless benzenediazonium hydroxide. The e-erystalline product bas the composition

and possesses all the properties of a true metallie salt; it is very soluble, and its solution is an S61; Claus, J. pr. Chem. 1894, [u], 50, 239; electrolyte. Moreover, this double salt couples Meyer and Jacobsen, Lehrhuch der Org. Chem. with β-naphthol and is converted by alkalis into the yellow syn-duzzocyanide. These properties correspond with those of the normal diszoninm salts, and the foregoing soluble oyanide is regarded as having the following constitution .

Hantzsch, 1901, 34, 4168).

The two pairs of diazocyanides from pchloraniline and p-anisidine were examined apectroscopically by Dobbie and Tinkler, who found that each pair gave almost identical ultre-violet absorption spectre, whereas the above soluble diazonium cvanide gave an entirely different spectrom. These results are in accordance with Hantzsch's view of the configuration of the diazo and diazonium cyanides (J.C.S. 1905, 87, 273).

It has, however, been suggested by several

VII. Diazosulphonates.

Although the stereochemical theory of the constitution of diszocyanides is supported by physical as well as chemical evidence, yet it is augnificant that the only other series of salts in which this isomerism has been detected is one derived from sulphurous acid, a substance resembling hydrocyanio scid in giving rise to organio isomerie derivativea which are atructur-

ally dissimilar.

The diazosulphonates, prepared by adding potassium sulphite to aqueous solutions of benzenoid diazonium chlorides, frequently exist in two differently coloured modifications, but in

most cases the syn. someride is too matable to be isolated in a pure state. Potassium syn-2:4-dilodobenzenedlazo-sulphonate (I) is an orango aubstance, whilst the anti- salt (II) is yellow:

The diszotised naphthylamines behave exceptionally, yielding only syn-diazosulphonates, which, on warming, pass, not into their antiseomerides, but into the corresponding azonaphthalenes (Hantzsch and Schmiedel, Ber. 1897, 30, 71).

The syn- and anti- modifications of potassium benzenediazosulphonate, C, H, N.N.SO, K, were found by Dobbie and Tinkler (i.e.) to have identical ultra-violet absorption spectra. fact supports the stereochemical theory of their structures (cf. Hantzsch, Ber. 1894, 27, 1726, 2099, 2586; Bamberger, shid. 2930, 3527; Bamberger, Ber. 1895, 28, 225, 444, 826; Pechmann, wid. 1902, is, 303).

VIII. Aliphatic Diazo- Compounds.

The amines of the aliphatic series, when treated with nitrous acid, generally loss their basic nitrogen, the amino group becoming replaced by hydroxyl. In some cases, bowever, the nitrogen is retained and an aliphatic diszo-(Hantzsch, Ber. 1900, 33, 2161; Euler and compound is produced, having the general

formula RHC | with the azo- group attached

entirely to the same carbon atom.

An alternative formulation has been auggested for the aliphatic diazo- compounds in which one nitrogen is quinquevalent, so that the general formula becomes :> CN:N (Thiele, Ber. 1911, 44, 2522; Forster and Cardwell, J.C.S. 1913, 103, 861 : ef. Ber. 1912, 45, 1654 ; 1916, 49, 1884.

Although glycine is converted by nitrous workers in this field that the isomerism of the acid or alkyl nitrites into glycollic acid, yet its ethyl ester gives rise to ethyl diazoacetate, N₂:CH·CC₂·C₂H₅, a yellow liquid freezing at -22° and hoiling at 143-144°/721 mm. (Curtius, Ber. 1883, 16, 2230).

One kg. of ethyl aminoacetate hydrochloride and 750 g. of sodium nitrite are added successively to 2 litres of water containing 5 g. of sodium acetate. The mixture is shaken until the temperature falls to 0°; 5 c.c. of 10% sulphurio acid and 500 c.c. of ether are added and the liquids thoroughly agitated. The ethereal layer is separated and the treatment with dilute acid and ether repeated until red fumes are evolved. The ethereal extracts are washed with dilute aqueous sodium carbonate till alkaline, dried with calcium chloride, and the solvent removed on the water hath. The yield of ethyl diazoacetate is 94.7% of the calculated quantity (Silberrad, J.C.S. 1902, 81, 600).

With iodine, ethyl diazoacetate yields diiodoacetic acid, and with aldehydes it con-

denses to form ketonic esters:

$$\begin{array}{c} \text{R-CHO} + \text{N}_2\text{:CH-CO}_2\text{-C}_2\text{H}_5 \\ \qquad \rightarrow \text{R-CO-CH}_2\text{-CO}_2\text{-C}_2\text{H}_5 \end{array}$$

One of the most remarkable reactions of this diazo- ester is its condensation with henzene and its homologues, giving rise to a series of esters containing seven-membered hydrocarbon rings (Buchner, Ber. 1896-1903, 29, 106; 30, 632, 1949; 31, 399, 402, 2004, 2241, 2247; 32, 705; 33, 184; 36, 3509; Annalen, 1908, 358, 1).

Ethyl diazoacetate when heated with alkalis undergoes a series of complex polymerio changes (Curtius, Ber. 1885, 18, 1283; 1906, 39, 1383, 3398, 4140; 1907, 40, 84, 815, 1176, 1194, 1470; 1908, 41, 3116, 3140, 3161; J. pr. Chem. 1888 [ii], 38, 408; Hantzsch, Ber. 1900, 33, 58; cf.Silberrad, J.C.S. 1902, 81, 598).

Diazo- derivatives are not obtained from free a-amino-carhoxylic acids, and only from those amino-esters of aliphatic acids containing the amino- group in the a-position with respect to the alkylated carboxyl group. The esters of those polypeptides which contain the aminogroup in the a-position to a carhimino group can also be transformed into diazo- derivatives.

The a-aminoketones (e.g. a-aminoacetophenone C_6H_5 : CO·CH₂·NH₂) give diazo-compounds (Angeli, Ber. 1904, 37, 2080), and so also do certain uric acid compounds, e.g. aminomethyl-

Diazomethane,
$$CH_2 < N$$
 or $CH_2: N : N$, the [B], 1122).

simplest aliphatic diazo- compound, is prepared by adding methyl-alcoholic potassium hydroxide to an ethereal solution of nitrosomethylurethane,

and distilling the mixture on the water-hath, when the distillate consists of an ethereal solution of the diazo- compound, the yield being 50% of the calculated quantity. This substance is also produced by the action of hydroxylamine on methyldichloramine:

Diazomethane is a poisonous yellow gas condensing to a yellow liquid boiling at -24° to -23° and solidifying in pale yellow crystals melting at -145°. It is a powerful methylating converting hydroxyl gronps into methoxyl, and alkylating primary and secondary hases. With iodine, diazomethane gives nitrogen and methylene iodide, and aldehydes are converted into methyl ketones (cf. Meyer, Monatsh. 1905, 26, 1295, 1311; Acree, Johnson, Brunel, Shadinger and Nirdlinger, Ber. 1908, 41, 3199; Schlotterheck, Ber. 1907, 40, 479; 1909, 42, 2559; Forster and Holmes, J.C.S. 1908, 93,

Diazomethane combines additively with unsaturated compounds; with acetylene it gives pyrazole (von Pechmann, Ber. 1898, 31, 2950), and with ethyl fumarate it furnishes ethyl pyrazolinecarhoxylate.

Diazomethane reacts with certain aldehydes, forming methyl ketones:

$$R \cdot CHO + CH_2N_2 = R \cdot CO \cdot CH_3 + N_2$$

There is an alternative reaction exhibited by those aldehydes which show a tendency to form hydrates. When such aldehydes react with diazomethane, substituted ethylene oxides are formed. p-Nitrohenzaldehyde gives a mixture of p-nitroacetophenone and p-nitrophenylethylene oxide,

Similarly chloral is converted into aaa-trichloropropyleno-βy-oxide, CCl3-CH-CH2

(Arndt and Eistert, Ber. 1928, 61, [B], 1118, with Amende, ibid. 1949).

A similar process occurs when acid chlorides treated with diazomethane yield either diazoketones or chloroketones. In the former alternative the acid chloride (I mol.) is added to an ethereal solution of diazomethane (2 mols.):

$$R \cdot COCI + CH_2N_2 = R \cdot CO \cdot CHN_2 + HCI$$

The hydrogen chloride liherated decomposes any excess of diazomethane; it also attacks the diazoketone so that the corresponding chloroketone may be prepared by the following reaction:

R·CO·CHN₂+HCl=R·CO·CH₂Cl+N₂

(Bradley and Rohinson, J.C.S. 1928, 1310, 1545, 2904; Arndt and Amende, Ber. 1928, 61,

Although for some time open-chain formulæ have been adopted for aliphatic diazo-compounds $R_2C=N \Rightarrow N$ and azides $R \cdot N=N \Rightarrow N$, there is no really convincing evidence for these formula. tions. An examination of the electron-diffraction pattern of the vapour of diazomethane suggests that the compound is in resonance

between the two phases $H_2C = \overline{N} = \overline{N}$ and H₂C-N≡N. Azomethane when similarly examined indicates the conventional formula $CH_3 \cdot N = N \cdot CH_3$, but with methyl groups in 600

65, 311). Determinations of the parachors of diazo-oxlde, C, H, CH, N.N OK, H, O. is certain axidos made by Lindomann and Thieles simulatly prepared from mitrosobemyjurethane. Ber. 1929, 61, [B], 1529) are in favour of the These motalite derivatives are very untable;

for aliphatic diazo- compounds R.C.

azides RN || is obtained from consideration of

the boiling points of many of these derivatives, which, if cyclic, should approximate to the boiling points of the corresponding halides, whereas an open chain atructure would require boiling points near those of the nitro-compounds. Actually the boiling points of the azides he near those of the bromides or between these and the boiling points of the iodides (cf. Forster and Newman, J.C.S. 1910, 97, 2572), whereas the boiling-points of the aliphatic diazo- compounds lie very near to those of the chlorides ; those of the nitro-compounds are always much higher (Sidgwick, J.C.S. 1929, 1108; Hantzsch, Ber. 1933, 66, [B], 1349).

Diazoethane, CH, CH.N, and phenyi-diazomethane, C, H, CH.N, have both been prepared (Hantzsch and Lehmann, Ber. 1902, 85, 897). The former closely resembles diano

methane, the latter is a dark red oil. Substitution increases considerably colour of these diazo- derivatives : dimethyl-

diazomethane, C(CH2)2 N2, is red and diphenyl diazomethane is obtained in bluish-red needles melting at 20° (Standinger, Ber. 1916, 49, 1884) Diszomethane should not be confused with

azomethane, CH, NiH-CH, the simplest azo-compound, which is produced by oxidising symmetrical dimethylhydrarine (hydrazome-thane) with chromic acid (Thiele, Ber. 1909, 42, 2575). Azomethane is a colourless gas condensing to a pale yellow liquid (b p. 1-5").
Diazomethanedisulphonic acid, a note-

worthy example of an aliphatic diazo-compound, results from the interaction of potassinm cyanide and potassium hisulphite in presence of caustic potash. The solution, acadified and treated with for method of preparation, see Zincke and nitrous acid, yields successively aminomethane-Tropp (Annalen, 1908, 363, 302). disulphonic acid and the diazo- compound. Sulphurous Acid and diagomethanedisulphonic acid yield an additive compound which, on boiling, furnishes hydrazine (von Pechmann, Ber. 1895, 28, 2374; 1896, 29, 2161).

Olazoacetone, CH, CO CH.N, a pale vellow liquid, is obtained from ammoscetylacetone by converting this base into diazoacetylacetone anhydride and treating this product with aqueous caustio alkali (Wolff, Aunalen, 1912, 394, 23).

Metaliic diazo- derivatives of the allphatic series are produced by treating mitrosoalkyl urethanes with concentrated caustic potash solution or ethereal potassium ethoxide. Potas- 71). sium methyidlazo-oxide,

CH.N:N:OK,H,O,

separates in white crystals, when nitrosomethyl. Br [CH2] Br .urethane is added to concentrated aqueous caustie potash at 0°. Polassium benzyl-

(Ref. 1925, 51, 15), 1529) are in avour to use annear measurement, as is also the evidence they are decomposed by water with explosive violence, yielding respectively diazomethans and phenyldazomethane (flantisch and Lehmann, Ber. 1902, 35, 897).

Bibliography. Hantzsch, "Die Diazoverbin-dungen, Ahren's Sammlung, Chemischer und

Chemisch-technischer Vorträge," 1902; Morgan, "Our Present Knowledge of Aromatic Diazo-Compounds," Brit. Assoc. Report, 1902 ; Eibner. "Zur Geschichte der Aromatischen Diazoverbindungen," 1903; Cain, "The Chemistry and Technology of the Diazo. Compounds," 1920; Saunders, "The Aromatic Diazo. Compounds and their Technical Applications," 1936

DIAZOOINITROPHENOL (4:6-dinitro-2-diazophenol). This example of the non metallic explosive primer has been patented by W. M. Dehn (U.S.P. 1404687), who claimed that it is suitable for general priming purposes, especially where fouling and amalgamation with metals are to be avoided. It is stated to be non-hygroscopic and can be stored under water at ordinary temperatures for months without undergoing any appreciable change. It is stable in contact with cold mineral acids but is decomposed by alkalis. Diazodinitrophenol can be safely and strongly compressed without its explosive power being impaired. When hested above 150°C., it explodes violently, but if finely divided and wetted with water it cannot be detenated by means of a No. 8 cap. Diazodinitrophenol is more powerful than fullmenate of mercury and when mixed with potassium chlorate in suitable proportions exhibits about 40% more energy on detonation than the most efficient mixtures of fulminate and chlorate.

DIAZOTYPE PAPER v. DIAZO. COX-POUNDS (this vol. p. 590a). 0-0'-DIBENZIL

PhCO CO-C₄H₄-C₅H₄-CO COPh,

OIBROMIN, 55 dibromodially barbiture

acid (p. BARBITURIC ACID, Vol. I, p. 623). OICARBOXYLIC ACIDS, Higher All-

phatie. There are three general synthetic processes for the production of these compounds : (i) The electrolysis of the potassium asks of the half-esters of lower dicarboxylic acids :

2CO,Et[CH,],CO,K -- CO, Et [CH,]21 CO, Et (Crum Brown and Walker, Anuslen, 1891, 261,

123; Walker and Lumsden, J.C.S. 1901, 79, 1200; Carmichael, ibid. 1922, 121, 2545; Fairweather, Proc. Roy. Soc. Edin. 1926, 48,

(u) Condensation of the appropriate dihalide with potassium cyanide, followed by hydrolysis :

→ CN [CH₂]_n CN CO.H[CH,],-CO.H

(iii) Condensation of the appropriate dihalide with ethyl sodiomalonate, followed by hydrolysis and decarboxylation, e.g.:

$$\begin{array}{c} \operatorname{Br} \cdot [\operatorname{CH}_2]_n \cdot \operatorname{Br} \\ \longrightarrow \quad (\operatorname{CO}_2 \operatorname{Et})_2 \cdot \operatorname{CH} \cdot [\operatorname{CH}_2]_n \cdot \operatorname{CH} (\operatorname{CO}_2 \operatorname{Et})_2 \\ \longrightarrow \quad \operatorname{CO}_2 \operatorname{H} \cdot [\operatorname{CH}_2]_{n+2} \cdot \operatorname{CO}_2 \operatorname{H} \end{array}$$

(Franke and Hankam, Monatsh. 1910, 31, 188; Chuit. Helv. Chim. Acta, 1926, 9, 264; 1927, 10. 167).

The dihalides necessary for procedures (ii) and (iii) may be obtained from a lower dibasic acid by the following series of operations:

$$\stackrel{\text{reduction}}{\longrightarrow} \mathsf{OH} \cdot \mathsf{CH}_2 \cdot [\mathsf{CH}_2]_n \cdot \mathsf{CH}_2 \cdot \mathsf{OH}$$

$$\downarrow \mathsf{HBr}$$

$$\mathsf{Br} \cdot [\mathsf{CH}_2]_{n+2} \cdot \mathsf{Br}$$

If desired, two more methylene groups may be introduced into the chain, thus:

(Chuit, *l.c.*).

Certain of these acids occur naturally; thus, (thansic tetradecane-1:14-dicarboxylic acid), CO₂H·[CH₂]₁₄·CO₂H, occurs, in combination, in the roots of Thapsia garganica L. bination, in the roots of Thapsia garganica L. (Canzoneri, Gazzetta, 1883, 13, 514) and as a glyceride in the wax of Juniperus Sabina (Bougault, Compt. rend. 1910, 150, 875), and heptadecane-1:17-, octadecane-1:18-, and nonadecane-1:19- dicarboxylic acids as glycerides in Japan wax (Geitel and van der Want, J. pr. Chem. 1900, [ii], 61, 153; Schaal, Ber. 1907, 40, 4785; Ruzicka, Stoll and Schinz, Helv. Chim. Acta, 1928, 11, 670).

The chief interest of these compounds lies.

The chief interest of these compounds lies, however, in their conversion into ketones by distillation of their thorium salts; the large-ring ketones so obtained are useful in perfumery as fixatives, cyclopentadecanone (exaltone) being particularly valuable in this respect. For a general review of this subject, see Ruzicka, Chem. and Ind. 1935, 54, 2.

The melting-points of the best-known straightchain higher dicarboxylic acids,

$$CO_2H\cdot[CH_2]n\cdot CO_2H$$
,

are given in the following table:

10 11 12 13 14 15 16 m.p.110° 128° 113° 126° 115° 124° 118° 125° 17 19 20 21 24 28 32 m.p. 119° 124° 118° 124° 123° 123°

H. N. R. DICENTRINE, C₂₀H₂₁O₄N, an alkaloid occurring in various Dicentra species, first is prepared by reducing the corresponding nitroisolated by Hcyl from D. formosa (0·1%) (Arch.

(von Braun, Ber. 1909, 42, 4550; von Braun Pharm. 1903, 241, 313), later by Asahina from D. and Danziger, ibid. 1912, 45, 1975).

pusilla (0·14-0·35%) and D. spectabilis (Arch. pusilla (0·14-0·35%) and D. spectabilis (Arch. Pharm. 1909, 247, 201) and by Manske from D. Pharm. 1909, 247, 201) and by Manske from *D. oregana* and *D. eximia* (0·31%) (Canad. J. Res. 1933, 8, 592; 1934, 10, 765). It crystallises from Et₂O,EtOH or CH₃·CO₂Et in prisms, m.p. 168–169°, [a]_D +62·1° (in chloroform). The salts are well crystallised, characteristic is dicentrine-methine, m.p. 158–159° (Manske *l.c.*; Ghose, Krishna, and Schlittler, Helv. Chim. Acta, 1934, 17, 919). Absorption spectrum, see Girardet (J.C.S. 1931, 2630) and Kitasato (A. 1927, 1094). Gadamer for theoretical reasons. 1927, 1094). Gadamer, for theoretical reasons, assigned the following constitution to dicentrine (Arch. Pharm. 1911, 249, 698)—

which was found to be correct by synthesis (Haworth, Perkin and Rankin, J.C.S. 1925, 2018; 1926, 29).

Osada (Amer. Chem. Abstr. 1928, 22, 3664) by splitting off from dicentrine the methylenedioxy- group with phloroglucinol-H2SO4 and completely methylating the phenolic base obtained glaucine. Dicentrine was obtained by methylating the phenolic alkaloid actinodaphnine (Ghose, Krishna, and Schlittler, l.c.). According to Iwakawa (Arch. exp. Path. Pharm. 1911, 64, 369) dicentrine in small

doses produces narcosis, in largo doses it causes convulsions, attacks the heart, and paralyses the respiratory centres.

Schl. DICHLORAMINE T, toluene-p-sulphondi-chloramide, C₆H₄Me·SO₂·NCl₂, is produced by treating toluene with chlorsulphonic acid, converting the sulphonic chloride thus obtained into the amide which is then chlorinated (see T. Aldoschin, G. Chimici, 1934, 28, 87). Under many trade names it is now of great importance in many fields, e.g. as a disinfectant (R. Dietzel, Arch. Pharm. 1928, 266, 123; D.R.P. 401011; Dan. P. 49704, etc.), as a bleaching agent in the textile industry (H. Tatu, Rev. gén. Teinture, 1933, 11, 883), as a disinfectant for agricultural purposes (U.S.P. 2002589), as an oxidising and chlorinating agent (A. Bradfield, J.C.S. 1928, 782; J. Koetschet, Helv. Chim. Acta, 1930, 13, 587; G. Schiemann, Z. angew. Chem. 1927, 40, 1032), etc.

2:5-DICHLOROANILINE.

Chim, 1868, [iv], 15, 259) with nickel and hydrogen (F.P. 621434) or hy chlorination and subsequent hydrolysis of 3 chloroacetanilide (Beilstein and Kurbatow, Annalen, 1879, 195, 215, 220). For preparation, see M. Kohn and

S. Fink, Monatsh. 1931, 58, 73).

2 5. Dichloroaniline has m p. 50°, h p. 250°. It undergoes the normal Skraup reaction to give 5.8 dichloroquinoline (F.P. 727528), hut is more valuable as the first component of a variety of azo. dyestuffs (G P. 112820, 193211, 251843, 256999; B.P. 329353, etc.), and others which, whilst not true azo- dyestuffa, yet empley 25dichlorodiazobenzene ealts as intermediates (G.P. 460087, B.P. 314899). It yields 25dichloroaniline-4-sulphonic acid on sulphonation with furning sulphuric acid at 170-180°, this is oxidised by potassium dichromate to 25-dichloroquinone (Noelting and Kopp, Ber. 1905, 38, 3513), and on diszotisation affords the diszoanhydride (Battegay, Ber. 1906, 39, 84), which yields azo- dyestuffe (G.P. 222991, 268599, 273934).

DICHROITE, a silicate of aluminium and magnesium with some ferrous iron replacing magnesium.

DICINCHONICINE (DICINCHO-NINE) v. CINCHONA ALKALOIDS, this vol.

DI-CITURIN. Trade name for the monepotassium salt of diacetyleitric seid (Pounes and Byrne, London), B.P.C. 1934.

DICKITE. A crystallised clay mineral. H.AlaSlaOa, differing slightly from keolinite in its optical characters and X-ray pattern. The minute crystals from Almwch in Anglesey were first described as kaolinite by A. B. Dick in 1888.

L, J. S. DICONQUININE v. CINCHONA ALKA-Lorps, this vol , p. 162a.

DICROTYL: ABS-octodiene, v. BUTADIENES AND POLYOLEFINS (non-conjugated).

DICTAMNINE, C11HeO1N, occurs in Dictamnus albus (white dittany; Fr. dictame blanc, Ger. Weisser Diptam), which belongs to the Rutacess. It was first isolated by Thoma (Ber. Dout. pharm. Ges. 1923, 33, 68; Arch. Pharm. 1930, 268, 39) (yield 0 3%), and later by Asahina, Ohta and Inubuse (Ber. 1930, 63, [B], 2045) from Skimmia repens Nakai (yield 0 2%). For extraction, see Asahins et al. (I.c.)

It forms colourless prisms, m.p. 132-133°, and is an optically inactive, uncaturated weak base, insol. H.O, col. warm EtOH and CHC!, lese in Et.O and CH. CO.Et; yields sparingly soluble chromate, picronolate m.p. 178°, chloro-aurate, chloroplatmate, and picrate m.p. 163°. The hydrochloride, m.p. 170° (decomp), hydrolyses in hot H₀O and the free base

An isomerio pseudo dictamnine was synthesised hy Asabina and Inuhuse (Ber. 1932, 65, [B], 61) For physiological action of dictamnine, see Thoma and Asahma.

Thoma (A. 1930, 383) further isolated from albus : dictamnolacione, C18H18O, mp 279-280°; frazinellone, C1.H1.O2, m.p. 117°, [a]n -38 39°; trigonelline and choline, dictamnio acid and dictamnal, see Acahina et al. (i.c.). Stimminging, C₁₄H₁₅O₄N, was isolated from the leaves of S. japonica Thunb. (yield 0 06%) hy Honda (Arch. exp. Path. Pharm. 1904, 52, 69). The close relationship to dictamine was suggested by the great similarity of the absorp-tion spectra. The alkaloid, a weak base, crystallases in prisms, mp. 176°, celts hydrolyse in H₈C, pierate, m.p. 195-197° (decomp.). 18 contains three OMe groups and, like dictamnine, is converted by methyliodide into isoskimmianine. Asahma and Inubuse (Ser. 1930, 63, [B] 2052) consider that ekimmisnine is 7-8-dimethoxydictamnine,

From Oriza japonica Terasaka isolated four alkaloids (Amer. Chem. Abstr. 1932, 26, 730; 1935, 29, 7337);

Orizine, CasHanDaN or CasHanDaN, [a]" +83.29°, m.p. 152 5° Kokuszgine, C, H,O,N, optically inactive,

mp. 201°. Kokusaginine, C₁₄H₁₃O₄N or C₁₄H₁₃O₄N,

m p 171°. Kokusaginoline, C₁₇H₁₁O₄N·13H₁O, mp. 283°.

The properties of these four new alkaloids are very esmalar to those of dictamnine and skimmiamino; the similarity also of the absorptionspectra is atriking. Probably all contain the same heterocyclic ring system. Terasaka supposee kokusagine to be 67-methylenedioxydictamnine. Schl.

OICTAMNOLACTONE v. DICTAMNINE. DICYANINE v. CYANINE DYES

OICYANODIAMIDE. (H,CN,), Obtained by polymeries cyanoguanidine. tion of cyanamide or, more conveniently, from opparates.

Commercial calcium cyanamide ("KalkstickWhen treated with methyliodide in a scaled stoff"). The polymerisation is facilitated by tube, dictamnine (1) is converted into aso-addition of small amounts of acid, insufficient dictamnine (II), which contains -NMe instead to combine with all the calcium present, and by of -OMe, a reaction characteristic of a and boding for a few hours. From the filtered y methoxyquinolnes. By oxidation and decar extract dicyanodiamide cryetallises in colouriess boxylation Asahina et al. (i.c.) finally obtained needles, m.p. 205°. On heating sodium cyan-2.4 dihydroxyquinoline, which makes the follow-ing formulæ probable:

cyanamide separates as an oil and on stirring or shaking polymerises in a few hours (Hermann, Monatsh. 1905, 26, 1025). The rate of polymerisation is dependent on the reaction of the solution used and is greatest at $p_{\rm H}$ 9·6, decreasing rapidly ahove and below this point (Buchanan and Barsky, J. Amer. Chem. Soc. 1930, 52, 195). On warming with dilute acid dicyanodiamide is converted into dicyanodiamidine (guanyl urea),

NH2(NH:)C·NH·CO·NH2,

which forms a characteristic insoluble compound with nickel, Ni(C₂H₅N₄O)₂ (Grossmann and Schück, Analyst, 1910, 35, 247; 1909, 34, 455; 1907, 32, 273, 394). This reaction has been utilised in the determination of nickel in the presence of cohalt (cobalt dicyanodiamidine is soluble in water). The nickel compound can be dried at 115-160° without decomposition. C. D. Garby (Ind. Eng. Chem. 1925, 17, 266) has adapted the reaction to the determination of dicyanodiamide, this is first converted to the -amidine, which is then precipitated as the nickel compound. Dicyanodiamide may also he determined as the insoluble compound with silver picrate (Johnson, J.S.C.I. 1921, 40, 125T). Reduction of dicyanodiamide with zinc and dilute hydrochloric acid yields guanidine (dicyanodiamidine probably heing formed intermedially (J. Bell, Sci. Proc. Roy. Dublin Soc. 1926, 18, 207)), and hydrocyanic acid which is rapidly reduced to the further stage of methylamine (Bamherger and Seeherger, Ber. 1893 1583). Dicyanodiamide, on treatment with a mixture of nitric and sulphuric acid is converted into nitrodicyanodiamidine, and heating with ammonium salts (chloride, nitrate, or thiocyanate) yields guanidine salts (Werner and Bell, J.C.S. 1920, 117, 1133; Ewan and Young, J.S.C.I. 1921, 40, 109), prohably hy intermediate formation of diguanide salts. Condensation with dialkyl cyanacetic or malonic esters produces pyrimidine derivatives, e.g. with ethyl diethyloyanacetate a condensation product is obtained which on treatment with sulphuric acid gives harhituric acid (Bayer & Co., G.P. 165223; Chem. Zentr. 1906, 1, 514). On autoclaving dicyanodiamide with liquid ammonia at 150° an 80% yield of, melamine, C₃N₃(NH₂)₂, is obtained. With ammonium carbonate, carbon dioxide, and water under similar conditions, 75% of melamine and 25% of guanidine carbonate are formed (W. Scholl et al., Ind. Eng. Chem. 1937, 29, 202). Condensation with formaldehyde in the presence of acid yields an artificial resin (F.P. 665210) which on addition of casein forms plastic moulding composition (B.P. 323047, 1927). The constitutional formula of dicyanodiamide has been variously described as NH₂(NH:)C·NH·C:N,(NH₂)₂C:N·C:N (Bamberger, Ber. 1883, 16, 1459; 1891, 24, 899; Pohl, J. pr. Chem. 1908, [ii], 77, 533; Prianischnikoff, J.S.C.I. 1909, 28, 724) and more recently as the Baumann formula:

F. Chastellain (Helv. Chim. Acta, 1935, 18, 1287) after a detailed study of the chain of reactions

 $\begin{array}{c} \text{acld} \\ \xrightarrow{} \text{dicyanodiamidine} \end{array}$

$$\begin{array}{ccc}
 & \text{NH} & \text{C:NH} \\
 & \text{(a)} & & \text{NH:C(NH}_2) \cdot \text{NH:CN} \\
 & & \text{(b)} & & & & & & & \\
\end{array}$$

and that in neutral and alkaline solutions and probably in the solid form it has the structure (a), whereas in acid solution this is transformed into (b).

The commercial fertiliser calcium cyanamide usually contains small amounts of dicyanodiamide which was formerly supposed to be the cause of seedling losses sometimes observed when the fertiliser was applied at or within a few days of sowing (Cowie, J. Agric. Sci. 1918, 9, 113). This is now known to he a fallacy (Lefort des Ylouses, Chim. et Ind. 1927, 18, 216). Moreover, the amount of dicyanodiamide in modern forms of calcium cyanamide is extremely small.

DIDIAL. Trade name for a preparation containing dial and ethylmorphine diallylharhiturate, administered as a powerful hypnotic (Ciba, London). B.P.C. 1934.

"DIENE SYNTHESIS" v. ALIZARIN AND ALLIED COLOURING MATTER, Vol. I, p. 208. m-DIETHYLAMINOPHENOL,

m.p. 78°, b.p. 201°/25 mm. is hest made by sulphonation of diethylaniline followed by alkali fusion of the *m*-sulphonic acid (D.R.P. 44792, U.S.P. 403678; for a review of other methods, see Sansome, Rev. Gén. Mat. Col. 1924, 28, 127).

m-Diethylaminophenol readily undergoes condensation with many compounds; 1-2-dicarboxylic acids (e.g. itaconic, citraconic, succinic, etc.) give rhodamine dyestuffs, Rhodamine B (Brilliant Rose B) being the derivative of phthalic acid. It is a source of triphenylmethane dyes (D.R.P. 205758, 229466), oxazine dyes (D.R.P. 300253), and many others, and is used as an anti-oxidant in ruhher technology (U.S.P. 1899120).

DIETHYLANILINE, C₆H₅·NEt₂, is produced by the classical methods of ethylation applied to aniline halogen hydrides; by treatment of aniline with diethyl sulphate in presence of lime (U.S.P. 1570203), with alcohol and sulphuric acid (Laptew, Anilin Farb. Ind. (Russ.), 1934, 4, 551), with ethyl chloride and lime under pressure (U.S.P. 1923697, 1994851), with ether under pressure (F.P. 768142), paraldehyde under pressure (F.P. 776613), etc. The simultaneous reduction and ethylation of nitro-

benzene with zino end ocetaldehyde (D.R.P. | vessel. The occurrence of diffusion in gases 491856) and meny cetalytic processes employ- thus becomes intelligible, end there is emple ing elcohol as the ethylating agent (Brown and experimental evidence for extending the kinetic Reid, J. Amer. Chem. Soc. 1924, 46, 1836; theory to the liquid state end for considering Roy, J. Indian Chem. Soc. 1928, 5, 383; F.P. that the molecules therein are elso in a state 669824) ero of perticular interest. A variety of methods employing chlorsulphonic seid (B.P. can, under certain conditions, change their 270300), phogene (B.P. 273923), phthabe acid position in the crystal lattice, and a large number (U.S.P. 1890246), maleto anhydride (U.S.P. of examples of diffusion through solids or no 1991787), etc., are aveilable for the removal of known. partielly alkylated by products. Diethylandine has m.p. -81°, h.p. 216 5°.

Diethylaniline is cheracterised by its quaternery eddition compound with benzyl chloride (m.p. 104°) (Morvel and co-workers, J. Amer. Chem. Soc. 1929, 51, 3638) and is recommended as a reagent for aluminium (Grosset, Ann. Soc. sci. Bruxelles, 1933, [B], 53, 16) and zme

(Eegriwe, Z. enal. Chem. 1928, 74, 225). Diethylanthne is used in the proparation of diphenylmethane derivatives (U.S.P. 1954484, 1803331) of azo dyestuffs and of dyestuffs belonging to the triphenylmethane and safran-ine series (F.P. 755338), etc. It also finds appli-cetion in the treetment of lubricating oils (F.P.

636332) and as an anti-oxident in rubber chemistry (D.R.P. 345160).

DIETZEITE. A double iodate and chromate of calcium crystallising in the monochine system, ducovered in 1891 in the sodium intrate deposits of Atacama, Chile, Analyses leed to the formula TCa(IO₃), 8CaCrO₂, but the sumple double salt formula Ca(IO₃), 2CaCrO₂ appears more probable. Distinct crystals are rare, the nuneral usually forming crystalline fibrous aggregates of a derk gold-yellow colour Spg. 7, 3698, It is soluble in hot weter; on cooling, crystals of hydrated colcum ichate (Ca(IO₃), 4H₃O₃) expansion, the sample calcum reduce Ca(IO₃), 4H₃O₄, colled by the colcum colours of the colours of the colcum colours of the colcum colours of the colours system, discovered in 1891 in the sodium natrate nodate Ca(103), called lautarite, occurs in the same deposits.

L. J. S. DIFFRACTAIC ACID, a lichen acid.

DIFFUSION. In any gescons mixture or liquid solution the composition ultimately becomes the same whatever the original distribu-

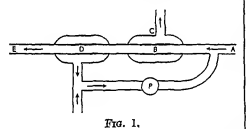
of constant motion. Even in solids the atoms

Diffusion of Gases.

The upwerd diffusion of heavy vepours, eq. bromme, into lighter gases has always been a popular lecture experiment, and in the latter part of the last century much work was done in investigating the quentitative laws that governed this process (see Loschmidt, Sitzungsgoverned this process (see Learning, Straing, St experimental difficulties ero met when direct diffusion experiments are ettempted, and it is much easier to study diffusion through porous pluge or through small apertures (effusion) In the above cases Graham ("Chemical and Physical Researches," T. and A. Constable, at the Edinburgh University Press, 1878, pp. 44, 88) was able to show that the volumes of different gases diffusing under standard conditions in a given time were inversely proportional to the given time were inversely proportional to the squeer protos of their densities. Knudsen (Ann. Physik, 1909, 28, 75) obtained the same result when gases were allowed to pass through a copulary tube (transpiration). The above relationship holds only approximately unless the mean free path of the gas molecules large compared with the aperture through which it is being driven—with a lerge hold the phenomenon is ohnously one of the viscous flow and not diffusion. The larger the above ratio beromet that better is Graham's law clowed feer Tumoffelf. the better is Graham's lew obeyed (see Timoffelf, Z. physikal. Chem. 1890, 6, 586; Donnen, Phil. tion of the gaseous or dissolved substances may Mag. 1900, 49, 423; Emich, Monetsh. 1903, 24, heve been. In very large systems, as, for 747; Knudsen, Ann. Physik, 1909, 28, 939). example, the atmosphere, there is a small [The study of diffusion in gases has been conchange in composition with height above the tinued by M. Trautz and co-workers as part of c eerth's surfece which is due to the change in general investigation into gaseous mixtures (see gravitational ettraction with the size of the Trantz and Müller, Ann. Physik, 1935, molecule. In the laboratory or factory this [v], 22, 313). Their work has hen entired effect is negligible except when the gravitational by J. Kuusmen (Ann. Physis, 1933, 171, 23, or eccelerating force becomes very large as in [445]. The problem of calculating diffusion the high-speed or ultra-centrifuge. Absential velocities in factory plant has been studied separations of the above type will not be considered in the present critical. The process by 401, Where equations audiable for the solution which the condition of equilibrium is reached is of practical problems are given. B. Kries and termed diffusion and it depends not on the N. Seljakov (Z. Physik, 1935, 94, 134) working circulation of the nnequally distributed snb- on mixtures of carhon dioxide end air have stence as e whole, but on the vibrations of studied the conditions of diffusion at high mindvidnal molecules. According to the knotte pressures. The ereporation and difficulties to the knotte pressures. The ereporation and difficulties the knotte pressures. The ereporation and difficulties the rest, is actually the scene of great activity as eatherst vapours into air has considerable on the part of the constituent molecules. These Eng. Chem. 1903, 26, 2631, 1003) has measured move with a high velocity and alter the direction of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of these processes for a multi-of of the velocity of the processes for a multi-of of the velocity of the processes for a multi-of of the velocity o of their motion only after they rousde with one liquids including weter, butyl alcohol, and another or with the walls of the containing toluene, and, to facilitate reference, the results

evaporation has been shown to consist of diffusion of the vapour from a supersurface layer into the gas phase and this diffusion is governed by the ordinary kinetic laws (E. Preston, Trans. Faraday Soc. 1933, 29, 1188; 1935, 31, 776, 1093). An ingenious apparatus for measuring the diffusion of vapours of volatile solids through gases is described by B. Topley and R. Whytlaw Gray (Phil. Mag. 1927, [vii], 4, 873). A small sphere of the substance under investigation is suspended from a fine quartz fibre which has been wound into a spiral, thus forming a spring balance which enables the loss in weight of the bead to be measured directly, the vapour being taken up by an adsorbent which is spread on the sides of the container. An accuracy of 2% was obtained for the passage of iodine vapour through air at temperatures between 14° and 30°C.

The principles of the diffusion of gases and vapours have important applications in the designing of pumps for high vacuum purposes. They have also been utilised by G. Hertz (Z. Physik, 1934, 91, 810; see also Barwich, ibid. 1936, 100, 166) in constructing an apparatus which enables the isotopes of the elements to be separated. One of the units from which the



apparatus was built up is shown diagram-matically in the fig. I. The gas enters the unit through a glass tube A in the direction shown by the arrow. It passes into a special porcelain diffusion tube B, which, by means of a grading glass, can be fused to the rest of the apparatus. A light fraction diffuses through B and is collected in the glass jacket C; it is pumped off and circulates with the gas in the previous unit. The remaining gas passes into another porous tube D through which a sample of medium density gas passes; the light fraction from the next unit is added to this and the mixture recirculated by means of a pump P. Finally the heavy remainder of the gas passes through E to the next unit where the whole cycle of operations is repeated. Thus in each unit there is a continuous process of separating the gas into three fractions, and by increasing the units it is possible to obtain any desired degree of separation. Herz has separated deuterium from hydrogen, also the neon and oxygen isotopes; indeed, once it is working, even small differences in density lead to effective separa-

DIFFUSION THROUGH METALS .- The passage of a gas across a diaphragm takes place not

are represented graphically. The process of only when the diaphragm is porous but also if it has the power of dissolving or absorbing the gas. For example, the ease with which hydrogen will pass through hot platinum or palladium sheet has been known for many years. recently further examples have been investigated and most of them have technical applications either in the wireless valve or metal industries. The more important of these papers are tabulated below.

N₂ through Cr.—G. Valensi, J. Chim. phys. 1929, 28, 152, 202.

H₂ through Cu.—E. O. Braaten and G. F. Clark, Proc. Roy. Soc. 1936, 153, 504; A. F. H. Ward, ibid. 1931, A, 133, 506, 522.

O2 through Cu.-F. Wilkins, ibid. 1930, A.

128, 407.

H₂O through Cu and N₂ through chrome on.—J. H. De Boer and J. D. Fast, Rec. trav. chim. 1935, 54, 970.

He through silica.—E. O. Braaten and G. F. Clark, J. Amer. Chem. Soc. 1935, 57, 2714;

Barrer, J.C.S. 1934, 387.

H2 through Ni, Cu, Mo and Fe; N2 through Mo.-C. J. Smithells and C. E. Ransley, Proc. Roy. Soc. 1935, A, 150, 172. H₂ through Al.—Idem, ibid. A, 1935, 152, 706.

CO through Ni.—Idem, ibid. A, 1936, 155,

195; A, 1936, 157, 292.

Ha through Pd.-V. Lombard and C. Eichner, Bull. Soc. chim. 1933, [iv], 53, 1176, ibid. 1935, [v], 2, 1555; Compt. Rend. 1935, 200, 1846: ibid. 1936, 202, 1777.

O₂, NH₃, S and P through Fe.—A. Bramley, F. W. Haywood, A. T. Cooper, and J. T. Watts,

Trans. Faraday Soc. 1935, 31, 707.

As a result of the above work the processes involved during the diffusion of gases through metals are known in outline at least. The first fundamental consideration is whether the gas passes through the crystals of the metal or penetrates through minute cracks and the filling at the crystal boundaries. Smithells and Ransley were able to show that the rates for the passage of hydrogen through fine and through large crystalline iron were identical and that diffusion must be through the body of the metal. On the other hand, Wilkins (l.c.) and Ward and Wilkins (Z. physikal, Chem. 1929, 144, 259) showed that in the case of oxygen through copper considerable lateral penetration round the boundaries occurs. A formula for the velocity of diffusion D across a metal plate from a pressure p on one side to zero pressure on the other was suggested by Richardson, Nicol, and Parnell (Phil. Mag. 1904, 8, 1), who gave

$$D = \frac{k}{d} \sqrt{pT} \cdot e^{-\frac{E}{RT}},$$

where k and E were constants, d was the thickness of the metal, T the absolute temperature, and R the gas constant. This equation ignores the effect of the surface of the metal in adsorbing the gas and it is now considered that unless this occurs the gas cannot penetrate the solid. Thus nitrogen will pass through iron or molybdenum but not through copper or nickel, while the inert gases will pass through glass and silica but not through metals (Barrer, I.c.; S. Bern-

¹ The quartz spring is usually spoken of as a McBain Microbalance.

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stem, Compt. rend. Acad. Sci. U.R.S.S. 1934, distribution of heat in thermal conductors. 1,230; H. Dunwald and C. Wagner, Z. physikal. Chem. 1034, B, 24, 53). Smithells and Raneley have shown that the adsorbed film must be allowed for before the Richardson equation would represent their experimental results. They used the following modification for the change of D with p at constant temperature:

$$D = \frac{l}{d} \sqrt{p} \left(\frac{ap}{1+\beta p} \right)$$

The values of the constants a and β agree with those obtained from direct adsorption measurementa and the necessity for adsorption to occur before diffusion is possible seems clearly established. It may be noted that the adsorp-tion process would more easily allow the molecules of gas to be aplit into atoms, and the relationship Dapi indicates that diffusion should occur in that state, Ward (i.c.) conadera the existence of atoms improbable, but B. Duhm (Z. Phyaik, 1935, 94, 434) has shown that for the case of hydrogen and palladium true dissociation into atoms can occur

In a recent communication J. Lynch (Nature, 1937, 40, 363) has suggested that the mechanical properties of metals may be considerably affected by the solution of gasee in them. In support of this contention be reported that the rigidity of a palladium wire was decreased by over 20% when it contained 450 times its volume of

discolved hydrogen.

Diffusion of Liquids and Dissolved Substances.

the solution to be diffused were carefully con. in experimenta lasting only 15 minutes. veyed to the bottom of the jar with the aid of a method has been carefully worked out by K. pipette. After a suitable time pertuens of 50 c.c. Sitte and can be used with very distressful warm of from the top; these Full details of the experimental precautions. portions evaporated to dryness, and the amount are given in the following series of papers and of aubstance that diffused into each layer was the complete apparatue is now marketed by thus ascertained. Graham's comparative experisous ascertained. uranam's comparative experial ments on the diffusive power of different above and the states in aqueous solution led him to draw a bid-stinction, which has become if the highest 29; R. Zuber and K. Sitte, ibid. 303, 300; importance, between the behaviour of easily Rivalk, 1932, 79, 275; R. Zuber, ibid. 290, 300; importance, between the behaviour of easily Rivalk, 1942, 79, 275; R. Zuber, ibid. 290, 300; importance, between the behaviour of easily Rivalk, 1942, 79, 275; R. Zuber, ibid. 290, 300; importance, between the behaviour of easily Rivalk, 1942, 79, 275; R. Zuber, ibid. 290, 300, 300; importance, between the behaviour of easily Rivalk, 1942, 79, 275; R. Zuber, ibid. 290, 300, 300; importance, between the short of easily state of the state cerned, the distinction in question is based on

The significance of the quantitative results nbtained by Graham was emphasised by Stefan $dS = -D q \cdot \frac{dc}{dt} \cdot dt$

Fick's Law is expressed by the formula

where dS is the amount of substance crossing a section of the diffusion column in time di. the sectional area of the column being q aq. cm. and deldx being the concentration gradient. The constant D is known as the diffusion co-officient. Stefan published tables from which the diffusion enefficient could be obtained directly from the analytical results after the solution had been divided into four portions. The method was modified by Oholm (Z. physikal. The method was modified by Obolm (Z. physika). Chem. 1904, 60, 309) and it is sometimes called after him. Its modern form is described by D. Krüger and H. Gruneky (bbd. 1930, 150, 115; 1934, 170, 161). A special sp-paratus for dividing the hapid into portion after diffusion was devised by E. Cohen and H. R. Brunn (bbd. 1925, 103, 349), who carried out some extremely careful deter-minations of the diffusion of symmetrical tetrabromoethane into tetrachloroethane be tween the temperaturee of 0 44° and 51.10°C. They discovered deviatione from Fick's Law amounting to about three times the magnitude of the possible experimental error. In any method the time required for diffusion to occur is considerable, and there is consequently ample opportunity for the solutions to be mixed by convection currents or by shaking, unless the apparatus is very firmly bedded and the temperature remains absolutely constant. These difficulties have been largely overcome by the The foundation of our knowledge of this introduction of a micro method by R. Eith subject was laid by the classical experiments (Physical Z. 1925, 28, 719). The cell is approximated from a firsh mir (*Chemical and Physical and Hayles (1818) in m. and is mounted on the Researches, T. & A. Constable, 1876, pp. 444—stage of a microscope through which the 600). In his experiments, 700 c. of water were progress of diffusion can be observed. Results the solution to be diffused were carefully compared to 1878, can be obtained the solution to be diffused were carefully compared to 1878, can be obtained to the control of the solution to be diffused were carefully compared to 1878, can be obtained to the control of the solution to be diffused where carefully compared to 1878, can be obtained to 1878.

efficients which is capable of giving very accurate results is to measure the amount of censes, the measure the amount of the following figures, representing approximate material that penetrates a fine eithered glass times of equal diffusion: bydrochloro acid 1, filter. The method was discovered by J. Hi. Northrop and M. L. Annon, (J. Gen. Phys. L. Morthrop and M. L. Annon, (J. Gen. Phys. Morthrop and M. L. Annon, (J. Gen. Phys. Morthrop and M. L. Annon, (J. Gen. Phys. L. Morthrop and M. L. Annon, (J. Gen. Phys. L. Morthrop and M. L. Annon, (J. Gen. Phys. L. Gen. Soc. 1931, 53, 69; 1933, 55, 432, 545). They have made numerous measurements with the apparatus and have used it in particular for (Sixtungsber, Akad, Wiss. Wien, 1878, 78, [61], the investigation of large molecules They 957; 1879, 79, [61], 161), who showed that they have abown that collouid particles can be made were in harmony with Fick's diffusion law, to diffuse normally and that the rate of which states that the diffusion of substances be used as a method of determining their molecular control of the state of the s in aqueous solution is comparable with the cular weight. Thus a value of 34,000 was obtained for egg albumin at its isoelectric point, and consequently the sharpness of the boundary which is in remarkable agreement with Sved-between the two liquids is not affected. berg's values of 34,500±1,000 by sedimentation equilibrium and 34,000 by osmosis. It is interesting to note that Svedberg could not obtain satisfactory values by diffusion, which McBain attributes to the use of buffer solutions, which he points out are difficult to maintain at identical concentrations on both sides of the diaphragm; secondly, they may cause a change in the size of the molecule under investigation, and thirdly, make analysis more difficult (see McBain, C. R. Dawson and H. A. Baker, J. Amer. Chem. Soc. 1934, 56, 1021). Contributions to the mathematics of the theory of diffusion have come from T. Katsurai and K. Kawashimo (Kolloid Z. 1936, 75, 37), also from W. G. Eversole and E. W. Doughty (J. Physical Chem. 1935, 39, 288), but the future development of the subject seems to call for the systematic tabulation of accurate

The difference in the rate of diffusion of heavy and light molecules through suitable membranes has been utilised for the removal of electrolytes from colloidal particles by dialysis. The membrane selected, however, is as far as possible semipermeable and consequently the process really belongs to the phenomena of osmosis.

Since the time of Graham the majority of experiments have confirmed that Fick's Law is at least approximately obeyed. A number of examples have, however, been found of materials that diffuse very much faster than anticipated and these substances were said to exhibit anomalous diffusion. This was first observed by R. O. Herzog and A. Polotzky with dyestuffs (Z. physikal. Chem. 1914, 87, 449); other examples, which were found, generally involved natural products, and explanations, like specific reaction with the solvent or lack of purity of the materials, were advanced. These had to be abandoned when H. Freundlich and D. Krüger showed that similar deviations could be obtained with substances of small molecular weight (Trans. Faraday Soc. 1935, 31, 906). For example, a solution of quinone diffusing into water obeys Fick's Law, but if both solutions are normal with respect to potassium sulphate anomalous results are obtained. The whole subject is critically reviewed by the above authors and they have shown that the anomalous results are attributable to the concentration gradient of the quinone causing the uniformly distributed substance (potassium sulphate) also to diffuse, owing to a mutual change in the solubility of the two solutes. In consequence, in the course of the diffusion process, a thin layer of the liquid on the border line of the two solutions becomes less dense than the solution immediately above. This inadmissible density gradient produces convection currents which obscure the normal process of diffusion. The correctness of this assumption was proved when care was taken that the sclution above

Diffusion of Solids

The possibility of diffusion processes occurring with solid materials was indicated by the manufacture of steel by the cementation process; but in spite of its importance industrially, it is only recently that much published work has appeared on the subject. The first example of solid diffusion to be studied was that of gold into lead; this was shown by Roberts-Austin (Phil. Trans. 1896, 187, 383) to occur to an appreciable extent in 40 days at temperatures between 100° and 200°C. In a subsequent paper (ibid. 1900, 67, 101) it was stated that the process could be detected at room temperatures but that time had to be measured in periods of years. A. E. van Arkel (Metall-Wirt. 1928, 7, 656) introduced an ingenious method of following diffusion when he coated a copper wire with a layer of nickel by electrolysis and was then able to detect any change by the variation of electrical conductivity of the composite wire. He found that the transfer of metal became appreciable above 800°C. O. Tanaka and Matano (Mem. Coll. Sci. Kyoto, 1931, 14, A, 59, 123) studied the temperature coefficient for the diffusion of silver into gold and of nickel into copper; and the same measurements were made for gold into lead by G. von Hevesy and W. Seith (Z. Elektrochem. 1931, 37, 528).
W. Seith and co-workers have in recent

years published a series of excellent experi-mental papers. Using a radioactive indicator thorium B-they followed its movement into crystals of lead and showed that in this symmetrical lattice the rate was practically the same in any direction (Z. Elektrochem. 1933, 89, 538). The experiments were then repeated with bismuth and it was found that the diffusion in a direction perpendicular to the crystalline axis was very much faster than when it was parallel to it. A technique was then worked out by which the course of diffusion could be followed by cutting thin layers of the metal and finding their composition by spectrum analysis. The rate of movement of magnesium, cadmium, nickel and mercury in lead; lead, mercury and platinum in cadmium; and platinum into zinc were then determined (*ibid*. 1934, 40, 318; Z. Metallk. 1932, 24, 193). The results obtained were then checked by measurements for a series of metals diffusing into silver at temperatures between 650° and 895°C. (Z. Elektrochem. 1936, 42, 570). The authors have been able to show that in general Fick's Law is obeyed and that diffusion D varies with temperature T according to the formula

$\log D = -A/T + B$

where A and B are constants. Similar conclusions were reached by A. Bramley and coworkers for the diffusion of carbon, sulphur, by the fact that the abnormality disappeared nitrogen and phosphorus into iron and steel (Trans. Faraday Soc. 1935, 31, 707). The effect was to a sufficiently marked degree less dense of a second element on the diffusion of a than the one below. It should be noted that the phenomenon occurs only in the boundary layers, results. The presence of 0.5% of carbon re608 DIFFUSION.

duced the velocity of the movement of sulphur reduced by hydrogen for 200 hours before by one half. On the other hand, the presence sitting, mutead of a minimum value being of sulphur or phosphorus almost entirely obtained, the process was nearly as fast as an prevents the movement of carbon and, in the the presence of the optimum amount of oxygen. Case of phosphorus, the carbon is driven before Amomalies of this type require, much expericase of prospriorus, the extron is driven scrops knowned of the trype require much experi it into the metal. The penetration of mitrogen mental work for their elicidetics, but, since is of particular importance owing to its applica-tions to the hardening of epecial sceles. It was an any commercial sample of steet, they may found that the presence of oxygen favoured the prove to be of the utmost importance. Intriding process, but when the specimens was

END OF THE TETED VOLUME